EXHIBIT 2

AO 88A (Rev. 12/20) Subpoena to Testify at a Deposition in a Civil Action

United States District Court

for the

| District of Delaware | | |
|--|--|--|
| Apple Inc. Plaintiff V. Masimo Corporation and Sound United, LLC Defendant) | Civil Action No. 22-cv-1378-MN-JLH | |
| SUBPOENA TO TESTIFY AT A DEF | OSITION IN A CIVIL ACTION | |
| To: Stryker Corp., 325 Corpoal c/o Registered Agent: U.S. Corporation Agents, Inc. | te Dr., Mahwah, NJ 07430 , 651 N. Broad St., Ste. 201, Middletown, DE 19709 | |
| (Name of person to whom | this subpoena is directed) | |
| Testimony: YOU ARE COMMANDED to appear at the deposition to be taken in this civil action. If you are an organizar party serving this subpoena about the following matters, or those or more officers, directors, or managing agents, or designate of these matters: SEE ATTACHMENT A | ation, you must promptly confer in good faith with the se set forth in an attachment, and you must designate one | |
| Place: Potter Anderson & Corroon LLP 1313 N. Market St., 6th Floor | Date and Time: | |
| Wilmington, DE 19801 | 08/11/2023 10:00 am | |
| The deposition will be recorded by this method: Ster | nographically, audiotaped, and videotaped | |
| Production: You, or your representatives, must also be electronically stored information, or objects, and must productial: See Schedule A (attached). Documents to be Bindu Palapura, Potter Anderson & Corroon bepalapura@potteranderson.com | permit inspection, copying, testing, or sampling of the | |
| The following provisions of Fed. R. Civ. P. 45 are attacked Rule 45(d), relating to your protection as a person subject to a series respond to this subpoena and the potential consequences of not | ubpoena; and Rule 45(e) and (g), relating to your duty to | |
| Date: 7/14/2023 | | |
| CLERK OF COURT | OR | |
| | /s/ Bindu A. Palapura | |
| Signature of Clerk or Deputy Clerk | Attorney's signature | |
| The name, address, e-mail address, and telephone number of the Plaintiff Apple Inc. | e attorney representing (name of party), who issues or requests this subpoena, are: | |
| Bindu Palapura 1313 N. Market St., Wilmington, DE 19801 302 | 2-984-6000 bpalapura@potteranderson.com | |

Notice to the person who issues or requests this subpoena

If this subpoena commands the production of documents, electronically stored information, or tangible things before trial, a notice and a copy of the subpoena must be served on each party in this case before it is served on the person to whom it is directed. Fed. R. Civ. P. 45(a)(4).

AO 88A (Rev. 12/20) Subpoena to Testify at a Deposition in a Civil Action (Page 2)

Civil Action No. 22-cv-1378-MN-JLH

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 45.)

| I received this subjon (date) | poena for (name of individual and title, if a | ny) | | |
|-------------------------------|--|--------------------------|------------------|------|
| ☐ I served the sub | poena by delivering a copy to the na | med individual as follow | /S: | |
| | | on (date) | ; or | |
| ☐ I returned the so | ubpoena unexecuted because: | | | |
| • | na was issued on behalf of the United ness the fees for one day's attendance | | • | |
| y fees are \$ | for travel and \$ | for services, for | or a total of \$ | 0.00 |
| I declare under per | nalty of perjury that this information | is true. | | |
| nte: | - | | | |
| | | Server's signa. | ture | |
| | | Printed name an | d title | |
| | | | | |
| | | Server's addr | ess | |

Additional information regarding attempted service, etc.:

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AO 88A (Rev. 12/20) Subpoena to Testify at a Deposition in a Civil Action (Page 3)

Federal Rule of Civil Procedure 45 (c), (d), (e), and (g) (Effective 12/1/13)

(c) Place of Compliance.

- (1) For a Trial, Hearing, or Deposition. A subpoena may command a person to attend a trial, hearing, or deposition only as follows:
- (A) within 100 miles of where the person resides, is employed, or regularly transacts business in person; or
- (B) within the state where the person resides, is employed, or regularly transacts business in person, if the person
 - (i) is a party or a party's officer; or
- (ii) is commanded to attend a trial and would not incur substantial expense.

(2) For Other Discovery. A subpoena may command:

- (A) production of documents, electronically stored information, or tangible things at a place within 100 miles of where the person resides, is employed, or regularly transacts business in person; and
 - (B) inspection of premises at the premises to be inspected.

(d) Protecting a Person Subject to a Subpoena; Enforcement.

(1) Avoiding Undue Burden or Expense; Sanctions. A party or attorney responsible for issuing and serving a subpoena must take reasonable steps to avoid imposing undue burden or expense on a person subject to the subpoena. The court for the district where compliance is required must enforce this duty and impose an appropriate sanction—which may include lost earnings and reasonable attorney's fees—on a party or attorney who fails to comply.

(2) Command to Produce Materials or Permit Inspection.

- (A) Appearance Not Required. A person commanded to produce documents, electronically stored information, or tangible things, or to permit the inspection of premises, need not appear in person at the place of production or inspection unless also commanded to appear for a deposition, hearing, or trial.
- **(B)** Objections. A person commanded to produce documents or tangible things or to permit inspection may serve on the party or attorney designated in the subpoena a written objection to inspecting, copying, testing, or sampling any or all of the materials or to inspecting the premises—or to producing electronically stored information in the form or forms requested. The objection must be served before the earlier of the time specified for compliance or 14 days after the subpoena is served. If an objection is made, the following rules apply:
- (i) At any time, on notice to the commanded person, the serving party may move the court for the district where compliance is required for an order compelling production or inspection.
- (ii) These acts may be required only as directed in the order, and the order must protect a person who is neither a party nor a party's officer from significant expense resulting from compliance.

(3) Quashing or Modifying a Subpoena.

- (A) When Required. On timely motion, the court for the district where compliance is required must quash or modify a subpoena that:
 - (i) fails to allow a reasonable time to comply;
- (ii) requires a person to comply beyond the geographical limits specified in Rule 45(c);
- (iii) requires disclosure of privileged or other protected matter, if no exception or waiver applies; or
 - (iv) subjects a person to undue burden.
- **(B)** When Permitted. To protect a person subject to or affected by a subpoena, the court for the district where compliance is required may, on motion, quash or modify the subpoena if it requires:

- (i) disclosing a trade secret or other confidential research, development, or commercial information; or
- (ii) disclosing an unretained expert's opinion or information that does not describe specific occurrences in dispute and results from the expert's study that was not requested by a party.
- (C) Specifying Conditions as an Alternative. In the circumstances described in Rule 45(d)(3)(B), the court may, instead of quashing or modifying a subpoena, order appearance or production under specified conditions if the serving party:
- (i) shows a substantial need for the testimony or material that cannot be otherwise met without undue hardship; and
 - (ii) ensures that the subpoenaed person will be reasonably compensated.

(e) Duties in Responding to a Subpoena.

- (1) Producing Documents or Electronically Stored Information. These procedures apply to producing documents or electronically stored information:
- (A) *Documents*. A person responding to a subpoena to produce documents must produce them as they are kept in the ordinary course of business or must organize and label them to correspond to the categories in the demand.
- **(B)** Form for Producing Electronically Stored Information Not Specified. If a subpoena does not specify a form for producing electronically stored information, the person responding must produce it in a form or forms in which it is ordinarily maintained or in a reasonably usable form or forms.
- (C) Electronically Stored Information Produced in Only One Form. The person responding need not produce the same electronically stored information in more than one form.
- **(D)** Inaccessible Electronically Stored Information. The person responding need not provide discovery of electronically stored information from sources that the person identifies as not reasonably accessible because of undue burden or cost. On motion to compel discovery or for a protective order, the person responding must show that the information is not reasonably accessible because of undue burden or cost. If that showing is made, the court may nonetheless order discovery from such sources if the requesting party shows good cause, considering the limitations of Rule 26(b)(2)(C). The court may specify conditions for the discovery.

(2) Claiming Privilege or Protection.

- (A) Information Withheld. A person withholding subpoenaed information under a claim that it is privileged or subject to protection as trial-preparation material must:
 - (i) expressly make the claim; and
- (ii) describe the nature of the withheld documents, communications, or tangible things in a manner that, without revealing information itself privileged or protected, will enable the parties to assess the claim.
- (B) Information Produced. If information produced in response to a subpoena is subject to a claim of privilege or of protection as trial-preparation material, the person making the claim may notify any party that received the information of the claim and the basis for it. After being notified, a party must promptly return, sequester, or destroy the specified information and any copies it has; must not use or disclose the information until the claim is resolved; must take reasonable steps to retrieve the information if the party disclosed it before being notified; and may promptly present the information under seal to the court for the district where compliance is required for a determination of the claim. The person who produced the information must preserve the information until the claim is resolved.

(g) Contempt.

The court for the district where compliance is required—and also, after a motion is transferred, the issuing court—may hold in contempt a person who, having been served, fails without adequate excuse to obey the subpoena or an order related to it.

For access to subpoena materials, see Fed. R. Civ. P. 45(a) Committee Note (2013).

ATTACHMENT A

SCHEDULE A

DEFINITIONS

The following terms shall have the meanings set forth below whenever used in any Definition, Instruction, Request for Production, or Deposition Topic.

- 1. As used herein, the terms "Stryker," "You," or "Your" means Stryker Corporation, Physio-Control Corporation, and all of their predecessors (merged, acquired, or otherwise), successors, subsidiaries, divisions, departments, and affiliates thereof, and all officers, directors, principals, agents, employees, attorneys, and other persons acting on their behalf.
- 2. As used herein, "Apple" means Apple Inc., all of its predecessors (merged, acquired, or otherwise), successors, subsidiaries, divisions, departments, and affiliates thereof, and all officers, directors, principals, agents, employees, attorneys, and other persons acting on their behalf.
- 3. As used herein, "Masimo" means Masimo Corporation, Cercacor Laboratories, Inc., and all their predecessors (merged, acquired, or otherwise), successors, subsidiaries, parents, sisters, divisions, departments, partnerships, and affiliates thereof, and all of their officers, directors, principals, agents, employees, independent contractors working under their control, attorneys, and other persons acting on their behalf.
- 4. As used herein, "Masimo Asserted Patents" means U.S. Patent No. 10,687,743 ("the '743 Patent"), U.S. Patent No. 10,722,159 ("the '159 Patent"), U.S. Patent No. 8,190,223 ("the '223 Patent"), U.S. Patent No. 10,736,507 ("the '507 Patent"), and U.S. Patent No. 10,984,911 ("the '911 Patent").
 - 5. As used herein, "Relevant Date" means September 20, 2012.
 - 6. As used herein, "Exhibits" means Exhibits 1 and 2 attached hereto.

- 7. As used herein, "Blood Oxygen and Heart Rate Features" means the product feature(s) relating to monitoring, measuring, sensing, detecting, and/or obtaining blood oxygen (SpO2) and/or heart rate measurements, including all hardware, software, firmware, components, modules, applications, and devices involved in such features, that were made or sold before the Relevant Date.
- 8. As used herein, "Product" means any machine, manufacture, apparatus, device, system, process, service, method, or instrumentality which is designed to function together electrically, mechanically, chemically, or otherwise, to achieve a particular function or purpose, including those offered for sale, sold, imported, or under development.
- 9. As used herein, "Relevant Products" means (1) the Physio-Control LIFEPAK 15 monitor/defibrillator, (2) Physio-Control CODE-STAT Data Review Software, (3) Physio-Control DT EXPRESS Data Transfer Software, (4) the products described in Exhibits 1 and 2, (5) any Product made or sold by or for You having Blood Oxygen and Heart Rate Features before the Relevant Date, (6) any related Products or modules having Blood Oxygen and Heart Rate Features before the Relevant Date, and (7) any other software programs, applications, or modules that display Blood Oxygen or Heart Rate from the Relevant Products.
- 10. As used herein, "Source Code" means any human-readable programming language or format that defines software, firmware or integrated circuits, including but not limited to, computer code, scripts, assembly, binaries, object code, Register Transfer Level ("RTL") descriptions, VHDL, Verilog, and other Hardware Description Language ("HDL") formats.
- 11. The term "Third Party" means any person or entity other than You, Masimo, or Apple.

- 12. As used herein, the term "document" shall have the full meaning ascribed to it by the Federal Rules of Civil Procedure and includes the original and every non-identical copy or reproduction in Your possession, custody, or control, and further is used in a broad sense to refer to any electronically stored information ("ESI") or any tangible object or thing that contains, conveys, or records information.
- 13. As used herein, the singular of any word shall include the plural, and the plural shall include the singular.
- 14. As used herein, "person" means any natural person or any business, legal, or governmental entity or association.
- 15. As used herein, "include" and "including" shall be construed to mean "without limitation," so as to give the broadest possible meaning to interrogatories and definitions containing those words.
- 16. As used herein, "and" and "or" shall be construed conjunctively and disjunctively so as to acquire the broadest meaning possible.
- 17. As used herein, "any" and "all" shall each be construed to mean "each and every," so as to acquire the broadest meaning possible.
- 18. As used herein, the singular of any word shall include the plural, and the plural shall include the singular.
- 19. As used herein, "related" or "relating" to any given subject means, without limitation, identifying, describing, discussing, concerning, assessing, stating, reflecting constituting, containing, embodying, tending to support or refute, or referring directly or indirectly to, in any way, the particular subject matter identified.

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- 20. As used herein, "identify" as applied to a document shall mean to specify: (a) the type of the document (i.e., whether it is a letter, memorandum, e-mail, etc.); (b) the document's title and general subject matter; (c) the number of pages of the document; (d) the date the document was prepared; (e) the name of each and every author, addressee, distributor, and recipient of the document; (f) the date each distributor distributed the document and the date each recipient received the document; and (g) the name of each person that has or had possession, custody, or control of the document.
- 21. Any term not specifically defined herein shall be defined in accordance with normal usage as well as with the Federal Rules of Civil Procedure and the Local Rules of the United States District Court for the District of Delaware.

INSTRUCTIONS FOR REQUESTS FOR PRODUCTION

- 1. Apple's Requests for Production seek responsive documents and information sufficient to answer each of the Requests that are known or available You or in Your possession, custody, or control. If, after exercising due diligence to secure the documents or information requested, You cannot fully respond to a Request for Production, state that such is the case and answer to the fullest extent possible, stating what responsive documents or information are available, what documents or information cannot be provided, why the documents or information are unavailable, and what efforts were made to obtain the unavailable documents or information. If documents or information responsive to a Request in this subpoena are in Your control, but not in Your possession or custody, promptly identify the entity with possession or custody.
- 2. Regardless of whether a production is in electronic or paper format, documents that were maintained together before production should be produced in the same form, sequence, organization, or other order or layout as they were maintained, including any labels, file folders, file jackets, covers, or containers in which such documents are located or with which such documents are associated. If copies of documents are produced in lieu of the originals, such copies should be legible and bound or stapled in the same manner as the original.
- 3. These Requests for Production shall be deemed continuing. Documents located, and information learned or acquired, at any time after Your response is due must be promptly supplemented at the place specified in this subpoena.
- 4. A copy of the Protective Order entered in this Action for the protection of any requested proprietary, confidential, or commercially sensitive information is attached hereto.

REQUESTS FOR PRODUCTION

- 1. Documents sufficient to identify and describe the functionality, features, and operation of the Blood Oxygen and Heart Rate Features of the Relevant Products and all components, modules, applications, hardware, software, and firmware contained therein, including, without limitation, user manuals, brochures, presentations, user guides, product literature, engineering specifications, circuit diagrams, architectural diagrams, bills of materials, technical manuals, product specifications, data sheets, laboratory notebooks, research papers, test data and results, analyses, invention disclosure forms, reports, service manuals, operator's manuals, implementation guides, white papers, product tutorials, and non-public documentation.
- 2. Documents sufficient to identify and describe the conception, design, research, development, testing, use, operation, maintenance, marketing, modifying, sale, offer for sale, and supply of the Relevant Products, including the persons and entities involved.
- 3. Documents, communications, and things comparing the Apple Watch to the Relevant Products.
 - 4. Other versions of the Exhibits and documentation related to the Exhibits.
- 5. Documents sufficient to show the earliest dates that each of the Relevant Products were first conceived; reduced to practice; and made, sold, used (including by third parties such as end users), offered for sale, in public use, and otherwise available to the public in the United States, including but not limited to documents relating to any conference, seminar, exhibition, convention, or trade show at which such Product is or was discussed, referred to, advertised, displayed, demonstrated, or shown, such as, without limitation, product specifications, catalogs, announcements, advertisements, brochures, articles, pamphlets, price lists, invoices, purchase orders, sales records, or other promotional, marketing, or sales materials.
 - 6. Publications related to the Relevant Products that were made available to the public.

- 7. Three samples of each Relevant Product.
- 8. Source Code sufficient to show the functionality of the Blood Oxygen and Heart Rate Features of the Relevant Products.
- 9. Documents sufficient to show the authorship and authenticity of all the documents produced in response to this subpoena.

DEPOSITION TOPICS

- 1. The functionality, features, and operation of the Blood Oxygen and Heart Rate Features of the Relevant Products.
- 2. The earliest dates that each Relevant Product was reduced to practice, made, sold, offered for sale, in public use, or otherwise available to the public.
- 3. The subject matter contained within the documents produced in response to Requests For Production herein, including the authentication thereof.
- 4. The authorship and authenticity of the documents produced in response to the Requests For Production herein.

EXHIBIT 1



LIFEPAK® 15 MONITOR/DEFIBRILLATOR

Operating Instructions



LIFEPAK® 15 MONITOR/DEFIBRILLATOR

Operating Instructions

Important Information

Device Registration

Please register your device at www.physio-control.com. This will ensure that you are notified of any product updates.

Text Conventions

Throughout these operating instructions, special text characters (for example, **CAPITAL LETTERS** such as **CHECK PATIENT** and **SPEED DIAL**) are used to indicate labels, screen messages, and voice prompts.

Version History

These operating instructions describe LIFEPAK 15 monitor/defibrillator devices with software revision 3313494-002 or later.

LIFEPAK, LIFEPAK CR, LIFEPAK EXPRESS, LIFENET, QUIK-LOOK, REDI-PAK, and QUIK-COMBO are registered trademarks of Physio-Control, Inc. CODE SUMMARY, Shock Advisory System, CODE-STAT, DT EXPRESS, SunVue, and cprMAX are trademarks of Physio-Control, Inc. Bluetooth is a registered trademark of Bluetooth SIG, Inc. Microstream, Smart CapnoLine, and FilterLine are registered trademarks of Oridion Systems Ltd. The Oridion medical capnography in this product is covered by one or more of the following US patents: 6,428,483; 6,997,880; 5,300,859; 6,437,316 and their foreign equivalents. Additional patent applications pending. PC Card is a trademark of the Personal Computer Memory Card International Association. Masimo, the Radical logo, Rainbow, and SET are registered trademarks of Masimo Corporation. Formula 409 is a registered trademark of The Clorox Company. Specifications are subject to change without notice.



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Chapter 1

Preface

This chapter provides a brief introduction to the LIFEPAK® 15 monitor/defibrillator and describes the product's intended use.

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Introduction

The LIFEPAK 15 monitor/defibrillator is a complete acute cardiac care response system designed for basic life support (BLS) and advanced life support (ALS) patient management protocols.

These operating instructions include information and procedures related to *all* features of the LIFEPAK 15 monitor/defibrillator. Your LIFEPAK 15 monitor/defibrillator may not have all of these features.

These operating instructions describe the operation of the LIFEPAK 15 monitor/defibrillator when the factory default settings are used. The factory default settings for all setup options are identified in table, Setup Options Factory Default Settings (on page 239). Your device may be set up with different default settings, based on your protocols. For information about changing default settings, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

IMPORTANT! Some LIFEPAK 15 monitor/defibrillator accessories are *not* interchangeable with accessories that are used with other LIFEPAK monitor/defibrillators. Specific accessory incompatibilities are noted in the related sections.

Intended Use

The LIFEPAK 15 monitor/defibrillator is intended for use by trained medical personnel. For information about training options, contact your local Physio-Control representative.

The LIFEPAK 15 monitor/defibrillator can be used in out-of-doors and indoor emergency care settings within the environmental conditions specified in Appendix A. The monitor/defibrillator is designed to be used during ground transportation except when specified otherwise.

Monitoring and therapy functions may only be used on one patient at a time. Manual mode monitoring and therapy functions are intended for use on adult and pediatric patients. Automated external defibrillation mode is intended for use on patients eight years of age and older.

For additional intended use information, and information about the indications and contraindications of the monitoring and therapy functions, see the individual sections identified below.

| • | ECG Monitoring | See Monitoring the ECG (on page 47) | Standard feature |
|---|---|--|------------------|
| • | 12-Lead Electrocardiography | See Acquiring a 12-Lead ECG (on page 58) | Optional |
| • | SpO ₂ , SpCO, and SpMet Monitoring | See Monitoring SpO2, SpCO, and SpMet (on page 68) | Optional |
| • | Noninvasive Blood Pressure Monitoring | See Monitoring Noninvasive Blood Pressure (on page 79) | Optional |
| • | End-Tidal CO ₂ Monitoring | See Monitoring ETCO2 (on page 86) | Optional |

Modes of Operation

| • | Invasive Pressure Monitoring | See Monitoring Invasive Pressure (on page 94) | Optional |
|---|-----------------------------------|---|------------------|
| • | Temperature Monitoring | See Monitoring Continuous Temperature (on page 103) | Optional |
| • | Vital Sign and ST Segment Trends | See Vital Sign and ST Segment Trends (on page 107) | Optional |
| • | Automated External Defibrillation | See Automated External Defibrillation (AED) (on page 119) | Standard feature |
| • | Manual Defibrillation | See Manual Defibrillation (on page 132) | Standard feature |
| • | Noninvasive Pacing | See Noninvasive Pacing (on page 141) | Standard feature |

Modes of Operation

The LIFEPAK 15 monitor/defibrillator has the following modes of operation:

- AED mode—for automated ECG analysis and a prompted treatment protocol for patients in cardiac arrest.
- **Manual mode**—for performing manual defibrillation, synchronized cardioversion, noninvasive pacing, and ECG and vital sign monitoring.
- Archive mode—for accessing stored patient information.
- **Setup mode**—for changing default settings of the operating functions. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.
- Demo mode—for simulated waveforms and trend graphs for demonstration purposes.
- **Service mode**—for authorized personnel to perform diagnostic tests and calibrations. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Service Manual*.

Chapter 2

Safety Information

This chapter provides important information to help you operate the LIFEPAK 15 monitor/defibrillator. Familiarize yourself with all of these terms and warnings.

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Terms

The following terms are used either in these operating instructions or on the LIFEPAK 15 monitor/defibrillator:

Danger: Immediate hazards that will result in serious personal injury or death.

Warning: Hazards or unsafe practices that may result in serious personal injury or death.

Caution: Hazards or unsafe practices that may result in minor personal injury, product damage, or property damage.

General Dangers and Warnings

The following are general danger and warning statements. Other specific warnings and cautions are provided as needed in other sections of these operating instructions.

| DANGER! | Explosion Hazard |
|---------|--|
| | Do not use this defibrillator in the presence of flammable gases or anesthetics. |
| WARNING | Shock Hazard |
| | The defibrillator delivers up to 360 joules of electrical energy. Unless properly used as described in these operating instructions, this electrical energy may cause serious injury or death. Do not attempt to operate this device unless thoroughly familiar with these operating instructions and the function of all controls, indicators, connectors, and accessories. |
| WARNING | Shock Hazard |
| | Do not disassemble the defibrillator. It contains no operator serviceable components and dangerous high voltages may be present. Contact authorized service personnel for repair. |
| WARNING | Possible Device Failure |
| | Do not modify the device. |

WARNING

Shock or Fire Hazard

Do not immerse any portion of this defibrillator in water or other fluids. Avoid spilling any fluids on defibrillator or accessories. Spilled liquids may cause the defibrillator and accessories to perform inaccurately or fail. Do not clean with ketones or other flammable agents. Do not autoclave or sterilize this defibrillator or accessories unless otherwise specified.

WARNING

Possible Fire

Use care when operating this device close to oxygen sources (such as bag-valve-mask devices or ventilator tubing). Turn off gas source or move source away from patient during defibrillation.

WARNING

Possible Electrical Interference With Device Performance

Equipment operating in close proximity may emit strong electromagnetic or radio frequency interference (RFI), which could affect the performance of this device. If use of equipment in close proximity is necessary, observe the device to verify normal operation in the configuration in which the device will be used. RFI may result in distorted ECG, incorrect ECG lead status, failure to detect a shockable rhythm, cessation of pacing, or incorrect vital sign measurements. Avoid operating the device near cauterizers, diathermy equipment, or other portable and mobile RF communications equipment. Do not rapidly key EMS radios on and off. Refer to Electromagnetic Compatibility Guidance (on page 261) for recommended distances of equipment. Contact Physio-Control Technical Support if assistance is required.

WARNING

Possible Electrical Interference

This defibrillator should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the defibrillator should be observed to verify normal operation in the configuration in which it will be used.

WARNING

Possible Electrical Interference

Using cables, electrodes, or accessories not specified for use with this defibrillator may result in increased emissions or immunity from electromagnetic or radio frequency interference (RFI) which could affect the performance of this defibrillator or of equipment in close proximity. Use only parts and accessories specified in these operating instructions.

WARNING

Possible Electrical Interference

This defibrillator may cause electromagnetic interference (EMI) especially during charge and energy transfers. EMI may affect the performance of equipment operating in close proximity. Verify the effects of defibrillator discharge on other equipment prior to using the defibrillator in an emergency situation, if possible.

WARNING

Possible Equipment Damage

Use only ECG cables that are specified for use with this device. Protection of the device against defibrillator discharge is dependent on the use of ECG cables that are specified by Physio-Control.

WARNING

Possible Improper Device Performance

Using other manufacturers' cables, electrodes, power adapters, or batteries may cause the device to perform improperly and may invalidate the safety agency certifications. Use only the accessories that are specified in these operating instructions.

WARNING

Possible Improper Device Performance

Changing factory default settings will change the behavior of the device. Changes to the default settings must only be made by authorized personnel.

WARNING

Possible Device Shutdown

Always have immediate access to a spare, fully charged, properly maintained battery. Replace the battery when the device displays a low battery warning.

WARNING

Safety Risk And Possible Equipment Damage

MR unsafe: Keep the defibrillator away from magnetic resonance imaging (MRI) equipment.

General Dangers and Warnings

WARNING

Possible Patient Burns

A defect in the neutral electrode connection on HF surgical equipment could cause burns at the lead or sensor site and damage to the monitor/defibrillator. Do not apply patient leads, sensors, or catheters when using high frequency (HF) electrosurgical equipment.

Note: The features of the LIFEPAK 15 monitor/defibrillator which could come in either direct or casual contact with the patient or caregiver during normal use are not made with natural rubber latex.

Chapter 3

Basic Orientation

This chapter provides a basic orientation to the LIFEPAK 15 monitor/defibrillator device and its controls, indicators, and connectors.

| Front View | 21 |
|-------------|----|
| Back View | 31 |
| Batteries | 32 |
| Home Screen | 34 |
| Alarms | 39 |
| Options | 41 |
| Events | 43 |

Front View

This figure shows the front of the LIFEPAK 15 monitor/defibrillator. The front of the device is described in the following sections.

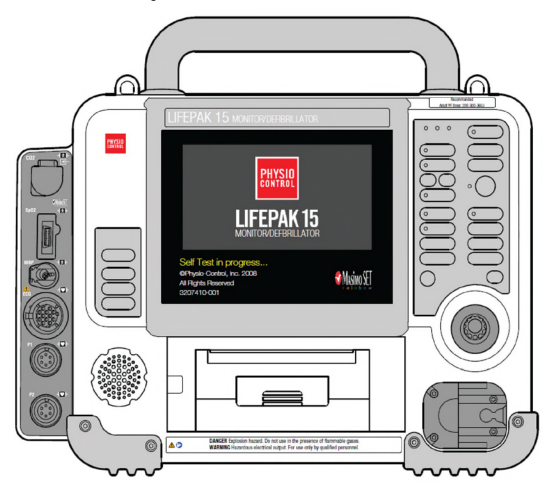


Figure 1 Front View

Area 1

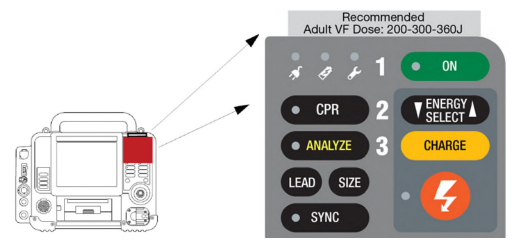


Figure 2 Area 1 Controls

| Table | 1 | Area | 1 | Controls |
|-------|---|------|---|----------|
| | | | | |

| able i Alea i Controls | | | | | |
|------------------------|---|---|--|--|--|
| CONTROL | DESCRIPTION | FOR MORE INFORMATION | | | |
| VF dose label | Physio-Control recommended energy dose for adult Ventricular Fibrillation (VF). | See <i>Biphasic Clinical Summaries</i> at www.physio-control.com/Biphasic | | | |
| ON | Turns device ON or OFF. LED illuminated when ON. Press and hold to turn device off. | | | | |
| ENERGY SELECT | Increases or decreases energy level in Manual mode. | See Manual Defibrillation (on page 132) | | | |
| CHARGE | Charges the defibrillator in Manual mode. | See Manual Defibrillation (on page 132) | | | |
| 4 | Shock button. Initiates discharge of defibrillator energy to patient. LED flashes when charging is complete. | See Manual Defibrillation (on page 132) | | | |
| 5 | Auxiliary power indicator. LED illuminated when defibrillator is connected to auxiliary AC or DC power source, whether defibrillator is turned on or off. | See Using the Power Adapter (on page 193) | | | |
| · · | Battery charging indicator. LED illuminated when installed batteries are fully charged. LED flashes when either battery is charging. LED is not illuminated when no batteries are installed or a battery is unable to be charged. | See AC Power Adapter Operation (on page 193) | | | |
| • * | Illuminated Service LED indicates a condition exists that prevents or could prevent normal defibrillator operation. | See General Troubleshooting Tips (on page 214) | | | |
| CPR | Controls CPR metronome. LED illuminated when metronome function is active. | See Using the CPR Metronome (on page 135) | | | |
| ANALYZE | Activates Shock Advisory System [™] (AED mode). LED illuminated when AED is analyzing the ECG, and flashes when user is prompted to push ANALYZE . | See Automated External Defibrillation (AED) (on page 119) | | | |
| LEAD | Changes ECG lead. | See Selecting ECG Lead (on page 48) | | | |
| SIZE | Changes ECG size. | See Changing ECG Size (on page 49) | | | |
| SYNC | Activates Synchronized mode. LED illuminated when Sync mode is active and flashes with detection of each QRS. | See Synchronized Cardioversion Procedure (on page 137) | | | |
| | CONTROL VF dose label ON ENERGY SELECT CHARGE CHARGE CPR ANALYZE LEAD SIZE | Physio-Control recommended energy dose for adult Ventricular Fibrillation (VF). Turns device ON or OFF. LED illuminated when ON. Press and hold to turn device off. ENERGY SELECT Increases or decreases energy level in Manual mode. CHARGE Charges the defibrillator in Manual mode. Shock button. Initiates discharge of defibrillator energy to patient. LED flashes when charging is complete. Auxiliary power indicator. LED illuminated when defibrillator is connected to auxiliary AC or DC power source, whether defibrillator is turned on or off. Battery charging indicator. LED illuminated when installed batteries are fully charged. LED flashes when either battery is charging. LED is not illuminated when no batteries are installed or a battery is unable to be charged. Illuminated Service LED indicates a condition exists that prevents or could prevent normal defibrillator operation. CPR Controls CPR metronome. LED illuminated when metronome function is active. Activates Shock Advisory System™ (AED mode). LED illuminated when AED is analyzing the ECG, and flashes when user is prompted to push ANALYZE. LEAD Changes ECG lead. SIZE Changes ECG size. Activates Synchronized mode. LED illuminated when Sync mode is active and | | | |

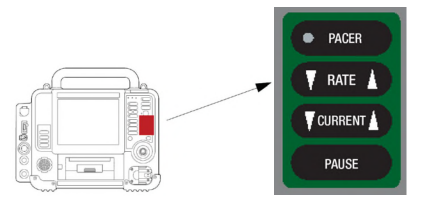


Figure 3 Area 2 Controls

Table 2 Area 2 Controls

| CONTROL | DESCRIPTION | FOR MORE INFORMATION |
|---------|---|--------------------------------------|
| PACER | Activates pacer function. LED illuminated when function is activated and flashes with each current pulse. | See Noninvasive Pacing (on page 141) |
| RATE | Increases or decreases pacing rate. | See Noninvasive Pacing (on page 141) |
| CURRENT | Increases or decreases pacing current. | See Noninvasive Pacing (on page 141) |
| PAUSE | Temporarily slows pacing rate. | See Noninvasive Pacing (on page 141) |

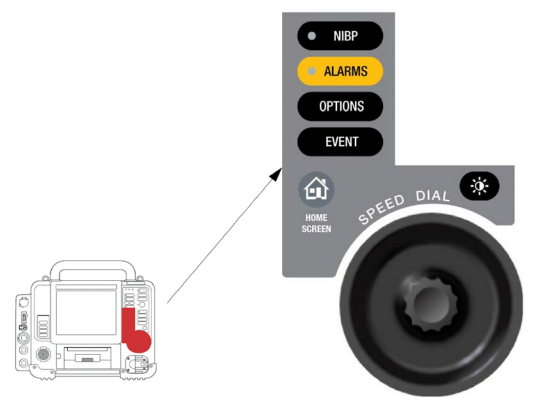


Figure 4 Area 3 Controls

Table 3 Area 3 Controls

| CONTROL | DESCRIPTION | FOR MORE INFORMATION |
|-------------|--|---|
| NIBP | Initiates blood pressure measurement. LED illuminated when BP measurement is being obtained. | See Monitoring Noninvasive Blood Pressure (on page 79) |
| ALARMS | Activates and silences alarms. LED illuminated when alarms are enabled and flashes when an alarm condition occurs. | See Alarms (on page 39) |
| OPTIONS | Accesses optional functions. | See Options (on page 41) |
| EVENT | Accesses user-defined events. | See Events (on page 43) |
| HOME SCREEN | Returns to Home Screen display. | See Home Screen (on page 34) |
| SPEED DIAL | Scrolls through and selects screen or menu items. | See Navigating the Home Screen (on page 37) |
| ∰ | Display mode button switches between color display and high contrast SunVue™ display. | |

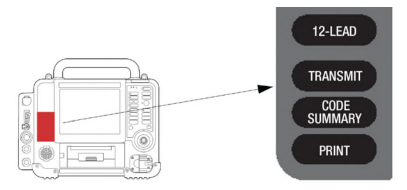


Figure 5 Area 4 Controls

Table 4 Area 4 Controls

| | *** | |
|--------------|---|--|
| CONTROL | DESCRIPTION | FOR MORE INFORMATION |
| 12-LEAD | Initiates acquisition of 12-lead ECG. | See Acquiring a 12-Lead ECG (on page 58) |
| TRANSMIT | Initiates transmission of patient data. | See Transmitting Reports (on page 183) |
| CODE SUMMARY | Prints CODE SUMMARY™ critical event record. | See CODE SUMMARY Report (on page 162) |
| PRINT | Starts and stops printer. | See How to Print a Current Report (on page 167) |

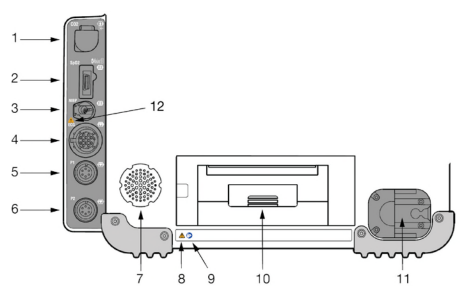


Figure 6 Area 5 Connectors, Speaker, and Printer

Table 5 Area 5 Connectors, Speaker, and Printer

| Table 5 / | riea o Connectors, | Speaker, and Finiter | |
|-----------|--------------------------|--|---|
| ITEM | LABEL | DESCRIPTION | FOR MORE INFORMATION |
| 1 | CO2 | FilterLine® set port | See Monitoring ETCO2 (on page 86) |
| 2 | SpO2/SpCO/ SpMet | Sensor cable port | See Monitoring SpO2, SpCO, and SpMet (on page 68) |
| 3 | NIBP | Pneumatic tubing port | See Monitoring Noninvasive Blood Pressure (on page 79) |
| 4 | ECG | Green electrically isolated ECG cable port | See Monitoring the ECG (on page 47) |
| 5 | P1 | Invasive pressure cable port | See Monitoring Invasive Pressure (on page 94) |
| 6 | P2 | Invasive pressure cable port | See Monitoring Invasive Pressure (on page 94) |
| 7 | Speaker | Projects device tones and voice prompts | |
| 8 | Symbol | General warning | See Warnings (on page 17) |
| 9 | Symbol | Follow instructions for use | |
| 10 | Printer | Door for 100 mm printer paper | See Loading Paper (on page 213) |
| 11 | Therapy cable receptacle | QUIK-COMBO® therapy cable and standard (hard) paddles cable receptacle | See Connecting and Disconnecting the Therapy Cable (on page 29) |
| 12 | Symbol | General warning | See Warnings (on page 17) |
| | | | |

Note: If your LIFEPAK 15 monitor/defibrillator is configured for temperature monitoring, P1 and P2 are replaced by a single port labeled TEMP. For more information about temperature monitoring, see Monitoring Continuous Temperature (on page 103).

Connectors

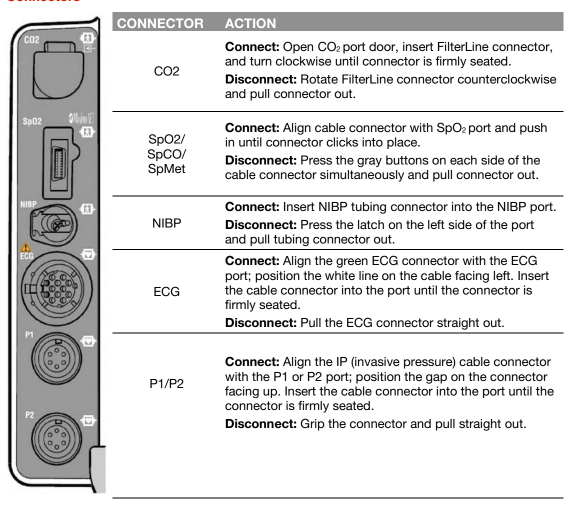


Figure 7 Connectors for IP Monitoring Configuration

Note: If your LIFEPAK 15 monitor/defibrillator is configured for temperature monitoring, P1 and P2 are replaced by a single port labeled TEMP. For more information, see the following figure, Connectors for Temperature Monitoring Configuration.

| | CONNECTOR | ACTION |
|-----------|-------------------------|--|
| C02 | CO2 | Connect: Open CO₂ port door, insert FilterLine connector, and turn clockwise until connector is firmly seated. Disconnect: Rotate FilterLine connector counterclockwise and pull connector out. |
| Sp02 | SpO2/ SpCO/ SpMet | Connect: Align cable connector with SpO ₂ port and push in until connector clicks into place. Disconnect: Press the gray buttons on each side of the cable connector simultaneously and pull connector out. |
| NIBP (FA) | NIBP | Connect: Insert NIBP tubing connector into the NIBP port. Disconnect: Press the latch on the left side of the port and pull tubing connector out. |
| | ECG | Connect: Align the green ECG connector with the ECG port; position the white line on the cable facing left. Insert the cable connector into the port until the connector is firmly seated. Disconnect: Pull the ECG connector straight out. |
| TEMPO | TEMP | Connect: Align the temperature adapter cable connector with the TEMP port. Insert the cable connector into the port until the connector is firmly seated. Disconnect: Grip the connector and pull straight out. |
| للم الم | | |

Figure 8 Connectors for Temperature Monitoring Configuration

Connecting and Disconnecting the Therapy Cable

WARNING

Possible Equipment Damage and Inability to Deliver Therapy

To help protect the therapy cable connector from damage or contamination, keep therapy cable connected to the defibrillator at all times. Inspect and test the therapy cable daily according to the Operator's Checklist in the back of this manual. Physio-Control recommends replacement of therapy cables every three years to reduce the possibility of failure during patient use.

IMPORTANT! The LIFEPAK 15 monitor/defibrillator QUIK-COMBO therapy cable and standard (hard) paddles have the same type of connector and connect to the defibrillator at the same location. These therapy cables are not compatible with other LIFEPAK defibrillator/monitors.

To connect a therapy cable to the defibrillator:

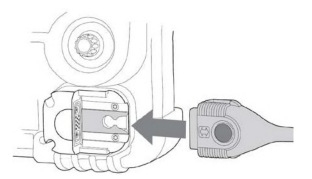


Figure 9 Connect Therapy Cable

- 1. Align the therapy cable connector with the receptacle.
- 2. Slide the therapy cable until you feel the connector lock in place. You will also hear a "click."

To disconnect the therapy cable from the defibrillator:

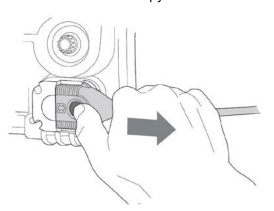


Figure 10 Disconnect Therapy Cable

- 1. Press the release button on the therapy cable connector.
- 2. Slide the therapy cable connector out.

Back View

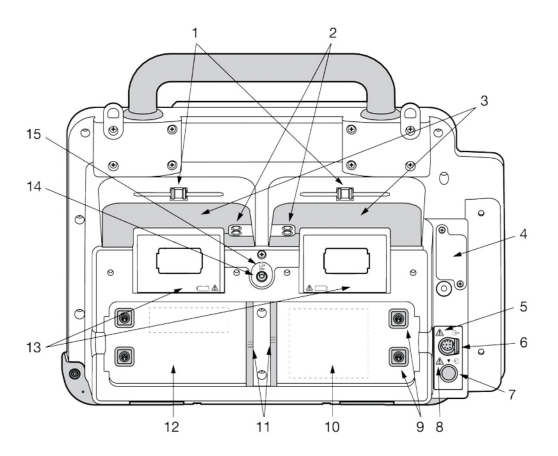


Figure 11 Back View

| FIG | FIGURE LEGEND | | | | |
|-----|---|----|---|--|--|
| 1 | Paddle retainers | 9 | Battery pins | | |
| 2 | Paddle test contacts | 10 | Battery well 1; includes serial number label | | |
| 3 | Standard paddle wells | 11 | Battery contacts | | |
| 4 | USB port cover | 12 | Battery well 2; includes Bluetooth label | | |
| 5 | See shock hazard warning (on page 32) | 13 | See battery warnings (on page 209) and stored battery warning (on page 210) | | |
| 6 | System connector | 14 | CO₂ exhaust port | | |
| 7 | Auxiliary power connector | 15 | See EtCO ₂ monitoring warnings (on page 87) | | |
| 8 | See power adapter warnings (on page 192) and battery warnings (on page 209) | | | | |

Back View

| Table 6 Back Vie | ew |
|------------------|----|
|------------------|----|

| LABEL | DESCRIPTION | FOR MORE INFORMATION |
|---|--|---|
| Battery wells, pins, and contacts | Each well holds one Lithium-ion battery. Two pins in each well transfer the battery power. Battery contacts transfer battery status information. See serial number label in battery well 1 for device part number, serial number, date of manufacture, and IP rating (dust and splash resistance). See Bluetooth label in battery well 2 for Bluetooth identification. See Using Bluetooth Wireless Communicationfor more information. | See Battery Maintenance (on page 209) |
| CO₂ exhaust port | Connects to a scavenger system when monitoring EtCO ₂ during use of anesthetics. | See Monitoring ETCO2 (on page 86) |
| Standard paddle wells, retainers, and test contacts | Paddle wells stow standard (hard) paddles. Retainers provide secure retention and quick removal of the paddles. Test contacts allow complete paddles defibrillation checks according to the Operator's Checklist. | See Standard Paddles (on page 154) and Operator's Checklist in the back of this manual |
| USB port cover | Protects USB port from the environment. | For future use |
| System connector | Connects device to a gateway or external computer for transfer of patient reports. Also provides real-time ECG output. | See Patient Records and Reports (on page 161) |
| Auxiliary power connector | Connects to an optional AC or DC power adapter. Allows use of auxiliary power source. | See Basic Orientation (on page 191) |

WARNING

Shock Hazard

All equipment connected to the system connector must be battery powered or electrically isolated from AC power according to IEC 60601-1. If in doubt, disconnect the patient from the defibrillator before using the system connector. Only use Physio-Control recommended data transmission cables. For more information, contact Physio-Control Technical Support.

Note: To prevent inadvertent depletion of the defibrillator batteries, disconnect external devices from the system connector when not in use.

Batteries

The LIFEPAK 15 monitor/defibrillator operates either on battery power using two Lithium-ion batteries, or with auxiliary power using the AC Power Adapter or DC Power Adapter. Batteries may be charged in the Station or Mobile Li-ion Battery Charger, the REDI-CHARGE™ Battery Charger, or in the monitor/defibrillator if it is connected to auxiliary power.

Note: Although the monitor/defibrillator can operate using auxiliary power with no batteries installed, at least one battery should be installed at all times. If the monitor/defibrillator loses power for more than 30 seconds, the device reverts to the user-configured default settings and begins a new patient record.

IMPORTANT! The LIFEPAK 15 monitor/defibrillator Lithium-ion batteries are not interchangeable with batteries that are used in other LIFEPAK defibrillators.

Routinely inspect batteries for damage or leakage. Recycle or discard damaged or leaking batteries.

Each battery has a fuel gauge that indicates the approximate charge level in the battery. Press the gray button above the battery symbol to check the battery's charge level prior to installing it in the defibrillator. The four battery indicators shown here represent approximate charge—greater than 70%, greater than 50%, greater than 25%, and 25% or less, respectively.



Figure 12 Battery Charge Indicators

Battery warning indicators are shown below. A single flashing LED indicates that the battery is very low and needs to be charged. Any two or more flashing LEDs indicate that the battery is faulty and should be returned to your authorized service personnel.



Figure 13 Battery Warning Indicators

Note: Older or heavily used batteries lose charge capacity. If a battery fuel gauge indicates fewer than four LEDs immediately after completing a charge cycle, the battery has reduced capacity. If the battery fuel gauge shows two or fewer LEDs after the battery completes a charge cycle, the battery should be replaced.

Batteries

To install a battery:

- 1. Confirm that the battery is fully charged, unless the battery will be charged in the monitor/defibrillator using the power adapter.
- 2. Inspect battery pins and contacts in the battery wells for signs of damage.
- 3. Align battery so battery clip is over the pins in the battery well.
- 4. Insert the end of the battery that is opposite the battery clip into the battery well.
- 5. Firmly press the clip end of the battery into the battery well until it clicks into place.
- 6. Repeat Step 1 through Step 5 to insert second battery.

To remove a battery, press the battery clip in and tilt the battery out of the battery well.

WARNING

Possible Loss of Power During Patient Care

Battery pins in the defibrillator may be damaged if batteries are dropped or forced into battery wells. Inspect pins routinely for signs of damage. Keep batteries installed at all times except when the device is removed from service for storage.

For information about battery maintenance, see Battery Maintenance (on page 209).

Home Screen

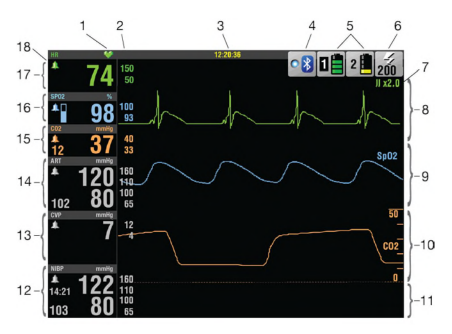


Figure 14 Home Screen

| FIG | FIGURE LEGEND | | | | |
|-----|-------------------|----|------------------------------|--|--|
| 1 | Heart symbol | 10 | Channel 3 | | |
| 2 | Alarm limits | 11 | Message area | | |
| 3 | Time | 12 | NIBP | | |
| 4 | Bluetooth icon | 13 | IP2 | | |
| 5 | Battery indicator | 14 | IP1 | | |
| 6 | Selected energy | 15 | EtCO ₂ | | |
| 7 | ECG Lead/Size | 16 | SpO ₂ /SpCO/SpMet | | |
| 8 | Channel 1 | 17 | Heart rate | | |
| 9 | Channel 2 | 18 | Alarm indicator | | |

The Home Screen is the main screen that displays ECG and other information. When a monitoring cable is attached to the device, the corresponding monitoring area on the screen is activated and the current patient values for that function are displayed. For example, when you connect an SpO₂ cable, the SpO₂ area is activated on the screen. SpO₂ values for the patient appear after the patient is connected. When the cable is disconnected, the SpO₂ patient values are replaced by dashes (--). Separate controls do not activate the monitoring functions, except for NIBP.

Each vital sign monitoring area is colored to match its waveform. This color scheme aids in associating the displayed waveform with its vital sign value. When a function does not have a waveform displayed, the vital sign area is gray.

WARNING

Failure to Detect a Change in ECG Rhythm

Heart rate meters may count internal pacing pulses during cardiac arrest or some arrhythmias. Do not rely entirely on heart rate meter alarms. Keep pacemaker patients under close surveillance.

Table 7 Home Screen

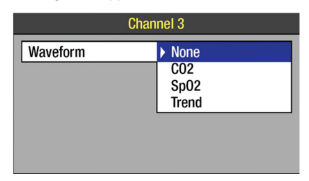
| Table 7 Home Screen | | FOR MORE INFORMATION |
|---|---|---|
| AREA Alarm limits | DESCRIPTION | |
| Alarm limits Limits display along the right side of the parameter. | | See Alarms (on page 39) |
| Heart symbol | Flashes with detected QRS signals. | |
| Alarm indicator | Indicates whether alarms are on or silenced. Absence of indicator means alarms are off. | See Alarms (on page 39) |
| Heart rate | Device accurately detects and displays heart rates between 20 and 300 beats per minute (bpm). If patient's heart rate is below 20 bpm or above 300 bpm, or pacing is active, dashes $()$ appear. If ECG is not active, the SpO ₂ or NIBP monitor can display pulse rate, indicated by PR (SPO ₂) or PR (NIBP). | |
| SpO2/SpCO/SpMet | Oxygen saturation level displays as a percentage from 50 to 100. Saturation below 50% displays as <50%. A fluctuating bar graph represents the pulse signal strength. When available and selected, the SpCO or SpMet value is displayed as a percent for 10 seconds, and then the SpO ₂ area reverts to the SpO ₂ reading. | See Monitoring SpO2, SpCO, and SpMet (on page 68) |
| EtCO2 | End-tidal CO₂ level displays in mmHg, Vol%, or kPa. Respiratory rate (RR) displays in breaths per minute. | See Monitoring ETCO2 (on page 86) |
| IP1/IP2 | Displays systolic, diastolic, and mean invasive pressures in mmHg. Two channels are available; default labels are P1 and P2. User-selectable labels include the following: • ART (arterial pressure) • PA (pulmonary artery pressure) • CVP (central venous pressure) • ICP (intracranial pressure) • LAP (left atrial pressure) | See Monitoring Invasive Pressure (on page 94) |
| Temp | Displays skin, esophageal, rectal, or bladder temperature. | See Monitoring Continuous Temperature (on page 103) |
| NIBP | Displays systolic, diastolic, and mean arterial pressures (MAP) in mmHg, and time to next BP, when interval is set. | See Monitoring Noninvasive Blood Pressure (on page 79) |

| AREA | DESCRIPTION | FOR MORE INFORMATION |
|-------------------|--|---|
| Time | Real or elapsed. | See LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device. |
| Bluetooth icon | Indicates <i>Bluetooth</i> capability. The LED is illuminated when a <i>Bluetooth</i> connection is established. Select this icon to access the <i>Bluetooth</i> setup menu. | See About Transmitting Patient Records and Reports (on page 175) |
| Battery indicator | Indicates presence of battery in battery well 1 and 2, relative level of charge, and battery in use. | See Battery Status Indicators (on page 38) |
| Selected energy | Selected defibrillation energy. | |
| ECG Lead/Size | Lead and size for ECG. | See Selecting ECG Lead (on page 48) |
| Channel 1 | Displays the primary ECG waveform and is always visible. | See Selecting ECG Lead (on page 48) |
| Channel 2 | Displays an additional waveform, a continuation of the Channel 1 ECG (cascading ECG), or a trend graph. | See The Pleth Waveform (on page 75) |
| Channel 3 | Displays an additional waveform or a trend graph. | See Displaying and Printing Trend Graphs (on page 111) |
| Message area | Displays up to two lines of status messages. | See Summary of Screen Messages (on page 245) |

Navigating the Home Screen

Use the **SPEED DIAL** to navigate around the Home Screen. As you rotate the **SPEED DIAL**, the individual vital sign areas and waveform channels on the Home Screen are outlined. If you outline a vital sign area or channel and then press the **SPEED DIAL**, a menu appears.

For example, rotate the **SPEED DIAL** to outline Channel 3, and then press the **SPEED DIAL**. The following menu appears.



- Rotate the SPEED DIAL to the desired setting.
- 2. Press the **SPEED DIAL** to select the setting.

Whenever a menu is displayed, the ECG is always visible in Channel 1. To return to the Home Screen from any menu, press the **HOME SCREEN** button.

Rotate and press the **SPEED DIAL** to select an option in a menu.

Battery Status Indicators

The Home Screen displays battery indicators that show the following information about the batteries installed in the defibrillator:

- Presence or absence of battery in battery well
- Battery in use
- · Battery charge state

IMPORTANT! Always check the battery charge level and ensure batteries are adequately charged before use.

When two batteries are installed, the defibrillator uses the battery with the lowest level of charge first. The battery in use is indicated by a white battery number in a black box. When a battery reaches the replace battery state, the defibrillator automatically switches to the other battery. When all battery capacity is exhausted, the defibrillator turns off. If you insert a charged battery and repower the device in less than 30 seconds, the defibrillator retains its settings. The following table provides a description of the various battery status indicators.

Table 8 Battery Status Indicators

| Table 5 Dattery | Otatus maicato | 10 |
|-----------------|--|--|
| INDICATOR | MEANING | DESCRIPTION |
| | Active battery | The defibrillator is using the battery in well 1 for power. Battery status indicators display up to four green bars. Each green bar represents approximately 25% remaining charge. For example, three green bars indicate about 75% remaining charge. |
| 1 | Low battery | Battery in well 1 is in use and is low. One yellow bar indicates 5% to 10% remaining charge. |
| | Very low battery | Battery in well 1 is in use and is very low. One red flashing bar indicates 0 to 5% remaining charge. The defibrillator automatically switches to the other battery only if adequate charge is available. If both batteries show red bars, the REPLACE BATTERY voice prompt occurs. |
| 2 ? | Unrecognized battery | Battery in well 2 is not in use. Battery communication failed or a non-Physio-Control battery is installed. The battery may power the defibrillator but the level of charge is unknown and low battery messages and prompts will not occur. |
| 1 | No battery installed or fault detected | No battery is installed in battery well 1, or a fault was detected in the battery in well 1 and the device will not use the battery. |

Note: When the defibrillator is operating on auxiliary power using a power adapter, the battery indicators show the battery charge level, but the well numbers are not highlighted. The **LOW BATTERY** and **REPLACE BATTERY** messages and prompts do not occur when operating on auxiliary power.

Note: Older or heavily used batteries lose charge capacity. If a fully charged battery is installed in the defibrillator and the battery status indicator shows less than four bars, the battery has reduced capacity. If a battery status indicator shows only one or two bars after a fully charged battery is installed, the battery has less than half the normal use time and should be recycled.

Alarms

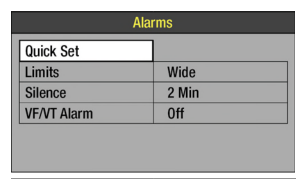
LIFEPAK 15 monitor/defibrillator alarms can be set up to be ON or OFF when the defibrillator is turned on. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

When alarms are set up to be ON, default limits are set. The limits temporarily appear to the right of the active vital signs. For all vital sign default alarm limits, see Alarm Limits (on page 237).

If alarms are set up to be OFF, press **ALARMS** to enable the alarms. Whether alarms are set up to be ON or are enabled by pressing **ALARMS**, they can only be turned off by pressing **ON** to turn off the device. If power is lost for less than 30 seconds, for example due to a system reset or changing the only active battery, alarm settings are restored automatically.

Setting Alarms

When you press **ALARMS**, the following menu appears:





Select **QUICK SET** to activate the alarms for all active monitoring functions.

The Quick Set limits automatically set high and low limits based on the patient's current vital sign values. For example, if the patient's HR is 70, selecting **WIDE** results in a high limit of 110 and a low limit of 45; selecting **NARROW** results in a high limit of 100 and a low limit of 50. The default is **WIDE**.

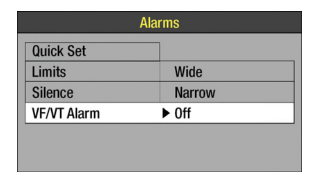
Select **LIMITS** to change alarm limits to **WIDE** or **NARROW**. See Alarm Limits (on page 237).

Select **SILENCE** to turn off the audible alarm for up to 15 minutes. If an alarm limit is exceeded while the alarm is silenced, the violated vital sign flashes and an alarm message appears, but the alarm tone remains silent.

If alarms are silenced for more than two minutes, an alert tone of two quick beeps sounds every 2.5 minutes. If alarms are silenced for two minutes, the alert tone sounds after 60 seconds.

Note: The heart rate display and corresponding heart rate alarm should not be relied upon to provide an indication of ventricular fibrillation. Turn on the VF/VT alarm.

Alarms



Select **VF/VT ALARM** to turn on continuous monitoring for ventricular fibrillation and ventricular tachycardia in Manual mode.

The VF/VT alarm indicator appears above the primary ECG when the alarm is ON.

Note: When the **VF/VT ALARM** is ON, you are limited to **PADDLES** lead or Lead **II** in Channel 1. See Selecting ECG Lead (on page 48).

Note: The VF/VT alarm is suspended when the metronome is active, the noninvasive pacemaker is on, or when standard paddles are attached and **PADDLES** lead is selected. The alarm is also suspended when the monitor/defibrillator is charging or is charged.

Managing Alarms

The alarm bell symbol indicates when alarms are ON or OFF. All alarms that are controlled by **QUICK SET** have equal priority. When alarms are ON and an alarm limit is exceeded, a tone sounds and the violated vital sign flashes.

To manage an alarm:

1. Press ALARMS. This silences the alarm for 2 minutes.

Note: After alarms are silenced by pressing the **ALARMS** button, an alert tone of two quick beeps sounds after 60 seconds.

- 2. Assess the cause of the alarm.
- 3. Assess the appropriateness of the limits settings (WIDE or NARROW).

If the patient is unstable, consider silencing the alarm for up to 15 minutes while attending to the patient. Do NOT reselect **QUICK SET**.



Possible Failure to Detect an Out of Range Condition

Reselecting **QUICK SET** resets the alarm limits around the patient's current vital sign values, which may be outside the safe range for the patient.

4. After the patient is stable, reselect QUICK SET, if necessary.

When alarms are ON, you can silence them preemptively for up to 15 minutes.

To silence alarms preemptively:

- 1. Press ALARMS.
- 2. Select SILENCE.
- 3. Select **SILENCE** duration of 2, 5, 10, or 15 minutes.

The message **ALARMS SILENCED** appears in the message area at the bottom of the Home Screen. If alarms are silenced for more than two minutes, an alert tone of two quick beeps sounds every 2.5 minutes. If alarms are silenced for two minutes, the alert tone sounds after 60 seconds.

Note: When you select **SILENCE**, the VF/VT alarm is not silenced.

Options

Press **OPTIONS** to display the Options menu. Rotate the **SPEED DIAL** to scroll through the choices. Press the **SPEED DIAL** to make a selection.

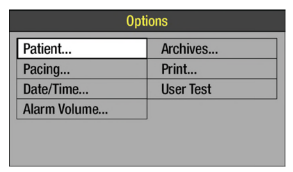
| Options | | |
|----------------|-----------|--|
| Patient | Archives | |
| Pacing | Print | |
| Date/Time | User Test | |
| Alarm Volume | | |
| | | |
| | | |
| | | |

Table 9 Options Menu Selections

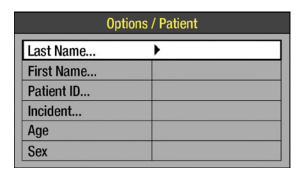
| SELECTION | DESCRIPTION | FOR MORE INFORMATION |
|--------------|---|---|
| Patient | Enter patient name, patient ID, incident, age, and sex. | See Entering Patient Data (on page 42) in next section |
| Pacing | Select demand or nondemand pacing. Set internal pacer detection ON or OFF. | See Noninvasive Pacing (on page 141) |
| Date/Time | Set date and time. Cycle power for change to take effect. | See LIFEPAK 15 Monitor/Defibrillator Setup Options for time display options. |
| Alarm Volume | Adjust volume for alarms, tones, voice prompts and CPR metronome. | |
| Archives | Access archived patient records. | See Managing Archived Patient Records (on page 169) |
| Print | Select report, format, mode, and speed for printing a current patient report. | See How to Print a Current Report (on page 167) |
| | | |

Entering Patient Data

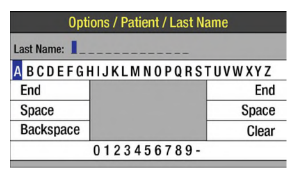
To enter patient data:



- 1 Press OPTIONS.
- 2 Use the SPEED DIAL to select PATIENT.



3 Select LAST NAME, FIRST NAME, PATIENT ID, INCIDENT, AGE, or SEX. (LAST NAME is selected in the example.)



- Rotate the **SPEED DIAL** to scroll through the characters and commands. Press the **SPEED DIAL** to make a selection. The selected character appears.
- 5 Repeat Step 4 until the name is complete.
- 6 Select END.

Three additional commands are available: **SPACE**—inserts blank space.

BACKSPACE—deletes last character and moves selection back one space.

CLEAR-clears all characters.

Events

Use the Events menu to annotate patient events. A selected event appears in the Event log of the CODE SUMMARY critical event record. Events can be customized in Setup mode. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

To select an event:

| Events | | |
|------------------|-------------|--|
| Generic | Intubation | |
| Oxygen | CPR | |
| IV Access | Epinephrine | |
| Nitroglycerin | Atropine | |
| Morphine | Lidocaine | |
| Cancel Last More | | |
| | | |

- 1. Press **EVENT** to display the Events menu.
- 2. Rotate the **SPEED DIAL** to scroll through the choices. Press the **SPEED DIAL** to make a selection.
- 3. Select **MORE** to display additional event selections.

Generic 12:20:30

When an event is selected, the event and time stamp appear in the message area on the Home Screen.

Notes:

- If you highlight an event but do not select it and the menu times out, a Generic event and time stamp are annotated in the event log.
- If you highlight an event but do not select it and then press **HOME SCREEN**, a Generic event and time stamp are annotated in the event log.
- Select CANCEL LAST to indicate that an incorrect event was selected. A Cancel Last event and time stamp print in the event log.

Chapter 4

Monitoring

This chapter describes the monitoring features of the LIFEPAK 15 monitor/defibrillator.

| Monitoring the ECG | 47 |
|---------------------------------------|-----|
| Acquiring a 12-Lead ECG | 58 |
| Monitoring SpO2, SpCO, and SpMet | 68 |
| Monitoring Noninvasive Blood Pressure | 79 |
| Monitoring ETCO2 | 87 |
| Monitoring Invasive Pressure | 94 |
| Monitoring Continuous Temperature | 103 |
| Vital Sign and ST Segment Trends | 107 |

Monitoring the ECG

Intended Use

The electrocardiogram (ECG) is a recording of the electrical activity of the heart. ECG monitoring allows for identification and interpretation of cardiac rhythms or dysrhythmias and calculation of heart rate. The ECG is obtained by placing either electrodes or paddles on the patient and allows the heart's electrical activity to be monitored and recorded.

ECG monitoring is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times; do not rely solely on the ECG monitor.

ECG Monitoring Warning

WARNING

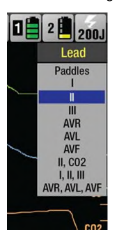
Possible Misinterpretation of ECG Data

The frequency response of the monitor screen is intended only for basic ECG rhythm identification; it does not provide the resolution required for diagnostic and ST segment interpretation. For diagnostic or ST segment interpretation, or to enhance internal pacemaker pulse visibility, attach the multi-lead ECG cable. Then print the ECG rhythm in diagnostic frequency response (DIAG) or obtain a 12-lead ECG.

Selecting ECG Lead

The LIFEPAK 15 monitor/defibrillator includes two methods for selecting or changing the ECG lead.

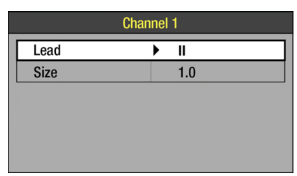
To select or change the displayed ECG lead using the **LEAD** button:



- Press LEAD. If any ECG lead currently appears on the Home Screen, the lead changes to PADDLES. If PADDLES lead is currently displayed, the lead changes to Lead II.
- 2. While the **LEAD** menu is displayed, press **LEAD** again or rotate the **SPEED DIAL** to the desired lead.

Note: If lead sets are predefined for Channels 2 and 3, the lead sets show on the menu. The ECG cable that is connected to the device, such as 3-lead or 5-wire, determines the leads you can select. For information about defining lead sets, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

To select or change the displayed ECG lead using the SPEED DIAL:



- 1. For the primary ECG, outline and select **CHANNEL 1** and then select **LEAD**.
- 2. Rotate the **SPEED DIAL** to the desired ECG lead.
- 3. Press the **SPEED DIAL** to select the ECG
- Repeat this procedure to select or change displayed ECG waveforms for Channels 2 and 3.

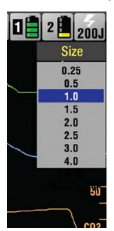
Note: The ECG shows dashed lines until the electrodes are connected to the patient.

Note: When the **VF/VT ALARM** is ON, you are limited to **PADDLES** lead or Lead **II** in Channel 1. See Setting Alarms (on page 39).

Changing ECG Size

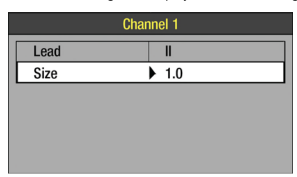
The LIFEPAK 15 monitor/defibrillator includes two methods for selecting or changing ECG size.

To select or change the displayed ECG size using the SIZE button:



- 1. Press SIZE.
- While the SIZE menu is displayed, press SIZE again or rotate the SPEED DIAL to the desired size.

To select or change the displayed ECG size using the SPEED DIAL:

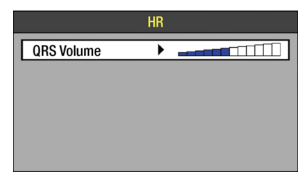


- 1. For the primary ECG, outline and select **CHANNEL 1** and then select **SIZE**.
- 2. Rotate the **SPEED DIAL** to the desired ECG size.
- 3. Press the **SPEED DIAL** to select the ECG size.

Adjusting the Systole Volume

To adjust the systole beep volume, use the **SPEED DIAL** to outline and select the **HR** area on the Home Screen.

The following menu appears:



- 1. Press the **SPEED DIAL** to select **QRS VOLUME**.
- Rotate the SPEED DIAL to the desired volume.
- 3. Press the **SPEED DIAL** to set the volume.

Note: The volume is reset to OFF each time the device is turned off.

Monitoring Using Paddle Accessories

To monitor ECG using paddles, you can use either QUIK-COMBO therapy electrodes or standard (hard) paddles. For more information about paddle accessories, see Paddle Accessory Options (on page 149).

Anterior-Lateral Placement

Anterior-lateral placement is the only placement that should be used for ECG monitoring using paddle accessories.

To place the therapy electrodes or paddles:

1. Place either the ♥ therapy electrode or APEX paddle lateral to the patient's left nipple in the midaxillary line, with the center of the electrode in the midaxillary line, if possible, as shown in the following figure.



QUIK-COMBO Therapy Electrodes

Standard Paddles

Figure 15 Anterior-Lateral Placement

2. Place the other therapy electrode or **STERNUM** paddle on the patient's upper right torso, lateral to the sternum and below the clavicle, as shown in the preceding figure.

Special Situations for Electrode or Paddle Placement

When placing therapy electrodes or standard paddles, be aware of the special requirements in the following possible situations:

Obese Patients or Patients with Large Breasts

Apply therapy electrodes or standard paddles to a flat area on the chest, if possible. If skin folds or breast tissue prevent good adhesion, it may be necessary to spread skin folds apart to create a flat surface.

Thin Patients

Follow the contour of the ribs and spaces when pressing the therapy electrodes or standard paddles onto the torso. This limits air spaces or gaps under the electrodes and promotes good skin contact.

Patients with Implanted Devices Such as Pacemakers or Defibrillators

If possible, place therapy electrodes or standard paddles away from implanted device.

Paddles ECG Monitoring Procedure

To monitor using standard paddles or therapy electrodes:

- 1. Press ON.
- 2. Prepare the patient's skin:
 - · Remove all clothing from the patient's chest.
 - Remove excessive chest hair as much as possible. Avoid nicking or cutting the skin if using a shaver or razor. If possible, avoid placing electrodes over broken skin.
 - Clean and dry the skin, if necessary. Remove any medication patches and ointment on the patient's chest.
 - Briskly wipe the skin dry with a towel or gauze. This mildly abrades the skin and removes oils, dirt, and other debris for better electrode adhesion to the skin.
 - Do not use alcohol, tincture of benzoin, or antiperspirant to prep the skin.
- 3. Apply the standard paddles or therapy electrodes in the anterior-lateral position. For therapy electrodes, confirm that the package is sealed and the Use By date is not passed. For standard paddles, apply conductive gel over the entire electrode surface.
- 4. Connect the therapy electrodes to the therapy cable.
- 5. Select PADDLES lead.

Monitoring Using ECG Cable Accessories

The following ECG cables, shown in the figure, are available for ECG monitoring with the LIFEPAK 15 monitor/defibrillator:

- 12-lead (either of 2 types)
- 3-lead
- 4-wire
- 5-wire

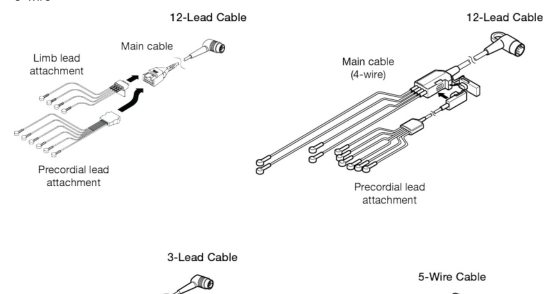
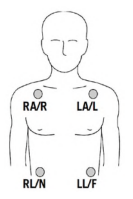


Figure 16 12-Lead, 3-Lead, 4-Wire, and 5-Wire ECG Cables

ECG Monitoring Procedure

To perform ECG monitoring:

- 1. Press ON.
- 2. Attach the ECG cable to the green connector on the monitor.
- 3. Identify the appropriate electrode sites on the patient as shown in the following figure.



| AHA Labels | | IEC La | IEC Labels | |
|------------|-----------|--------|------------|--|
| RA | Right Arm | R | Right | |
| LA | Left Arm | L | Left | |
| *RL | Right Leg | N | Negative | |
| LL | Left Leg | F | Foot | |

^{*}Note: Not used for 3-lead cable.

Figure 17 Limb Lead Electrode Placement

- 4. Prepare the patient's skin for electrode application:
 - Shave excessive hair at electrode site.
 - For oily skin, clean skin with alcohol pad.
 - Gently scrape skin to remove surface layer of dead cells and improve conduction of electrical signals.
 - Avoid locating electrodes over tendons and major muscle masses.
 - Clean and dry the skin.
- 5. Apply ECG electrodes:
 - Confirm that the package is sealed and the Use By date is not passed.
 - · Attach an electrode to each of the lead wires.
 - Grasp electrode tab and peel electrode from carrier.
 - Inspect electrode gel and make sure gel is intact (discard electrode if gel is not intact).
 - Hold electrode taut with both hands. Apply the electrode flat to the skin. Smooth tape outwardly. Avoid pressing the center of the electrode.
 - Secure the trunk cable clasp to the patient's clothing.

Note: Ensure the electrodes do not contact any other conductive parts, including earth (ground).

Note: Electrode quality is critical for obtaining an undistorted ECG signal. Always check the date code on electrode packages for expiration date before using on a patient. Do not use electrodes that have expired. Disposable electrodes are intended for a single use.

6. Select the desired ECG lead on the monitor screen.

Monitoring the ECG

- 7. If necessary, adjust ECG size for accurate heart rate counting.
- 8. Press **PRINT** to obtain an ECG printout.

Precordial Lead ECG Monitoring

The precordial (chest) leads (see Table, ECG Leads Color Codes (on page 54)) can be used for monitoring when using the 12-lead cable or 5-wire cable.

To perform precordial lead ECG monitoring:

- 1. Insert the precordial lead attachment into the main cable as shown in Figure, 12-Lead, 3-Lead, 4-Wire, and 5-Wire ECG Cables (on page 52).
- 2. Place the precordial lead electrodes on the chest as described in the 12-lead ECG procedure and shown in Figure, Precordial Lead Electrode Placement (on page 60).

Note: When using a 5-wire cable, attach the limb leads as described in ECG Monitoring Procedure (on page 53), and place the C-lead electrode on the chest in the precordial position desired. Note that the LIFEPAK 15 monitor labels the ECG for this lead as V1 on the screen and printout, regardless of the location of the C-lead electrode.

Leads Off

If an electrode or lead wire disconnects during ECG monitoring, the monitor emits an audible alarm and displays a **LEADS OFF** message. The ECG trace becomes a dashed line. The alarm and messages continue until one of the following actions is performed:

- · The lead wire is reconnected
- The lead selection is changed to a lead using connected lead wires
- Power is cycled.

Color Coding for ECG Leads

The lead wires and the electrode snaps for the patient ECG cable are color coded according to American Heart Association (AHA) or International Electrotechnical Commission (IEC) standards as listed in the following table.

Table 10 ECG Leads Color Codes

| LEADS | AHA LABEL | AHA COLOR | IEC LABEL | IEC COLOR |
|------------------|-----------|-----------|-----------|-----------|
| Limb Leads | RA | White | R | Red |
| | LA | Black | L | Yellow |
| | RL | Green | N | Black |
| | LL | Red | F | Green |
| | С | Brown | С | Brown |
| Precordial Leads | V1 | Red | C1 | Red |
| | V2 | Yellow | C2 | Yellow |
| | V3 | Green | C3 | Green |
| | V4 | Blue | C4 | Brown |
| | V5 | Orange | C5 | Black |
| | V6 | Violet | C6 | Violet |

Chapter 4 | Monitoring

Monitoring Patients Who Have Internal Pacemakers

The LIFEPAK 15 monitor/defibrillator internal pacemaker detection feature can be used to help identify internal pacemaker pulses on the printed ECG. When enabled, this feature uses lead V4 to detect internal pacemaker pulses. If V4 is not available because it is not attached or is too noisy, Lead II or Paddles Lead is used.

When the internal pacemaker detection feature is ON, the LIFEPAK 15 monitor/defibrillator annotates a hollow arrow on the printed ECG if internal pacemaker pulses are detected. Patient history and other ECG waveform data, such as wide QRS complexes, should be used to verify the presence of an internal pacemaker. False annotations of this arrow may occur if ECG artifacts mimic internal pacemaker pulses. If false annotations occur frequently, deactivate the detection feature using the **OPTIONS / PACING / INTERNAL PACER** menu (see Options (on page 41)).

The LIFEPAK 15 monitor/defibrillator typically does not use internal pacemaker pulses to calculate the heart rate. However, when using therapy electrodes or standard paddles to monitor in **PADDLES** lead, the monitor may detect internal pacemaker pulses as QRS complexes, resulting in an inaccurate heart rate.

Large amplitude pacemaker pulses may overload the QRS complex detector circuitry so that no paced QRS complexes are counted. To help minimize ECG pickup of large unipolar pacemaker pulses, place ECG electrodes so the line between the positive and negative electrodes is perpendicular to the line between the pacemaker generator and the heart.

Smaller amplitude internal pacemaker pulses may not be distinguished clearly in **PADDLES** lead. For improved detection and visibility of internal pacemaker pulses, turn on the internal pacemaker detector function using the **OPTIONS / PACING / INTERNAL PACER** menu or connect the ECG cable, select an ECG lead, and print the ECG in diagnostic frequency response. For information about configuring internal pacemaker detection, see the Pacing Setup menu in the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

Troubleshooting Tips

If problems occur while monitoring the ECG, check the table below for aid in troubleshooting. For basic troubleshooting problems, such as no power, see General Troubleshooting Tips (on page 214).

Table 11 Troubleshooting Tips for ECG Monitoring

| Table 11 Troubleshooting Tip | | |
|---|--|---|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
| Any of these messages displayed: | Therapy electrodes not connected | Connect therapy electrode. |
| CONNECT ELECTRODES CONNECT ECG LEADS ECG LEADS OFF XX LEADS OFF | One or more ECG electrodes disconnected | Connect ECG electrode. |
| | ECG cable is not connected to monitor | Connect ECG cable. |
| | Poor electrode-skin contact | Reposition cable or lead wires to prevent electrodes from pulling away from patient. |
| | | Secure trunk cable clasp to patient's clothing. |
| | | Prepare skin and apply new electrodes. |
| | PACER was pressed. The monitor automatically switched to Lead II, but ECG leads are not connected. | Connect ECG leads and initiate pacing. |
| | Broken ECG cable lead wire | Select another lead. |
| | | Select PADDLES lead, and use standard paddles or therapy electrodes for ECG monitoring. |
| | | Check ECG cable continuity. |
| Screen blank and ON LED | Screen not functioning | Print ECG on recorder as backup. |
| illuminated | properly | Contact service personnel for repair. |
| Systole beeps not heard or | Volume too low | Adjust volume. |
| do not occur with each QRS complex | QRS amplitude too small to detect | Adjust ECG size. |
| Displayed heart rate (HR) different than pulse rate | ECG size set too high or too low | Adjust ECG size up or down. |
| | Monitor detecting the patient's internal pacemaker pulses | Change monitor lead to reduce internal pacemaker pulse size. |
| Displayed heart rate (HR) different from displayed | ECG size set too high or too low | Adjust ECG size up or down. |
| ECG waveform | Monitor detecting the patient's internal pacemaker pulses | Change monitor lead to reduce internal pacemaker pulse size. |
| | | |

Chapter 4 | Monitoring

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|---|---|
| Monitor displays dashes () instead of heart rate | Heart rate is < 20 bpm | Use ECG printout to calculate heart rate. |
| | Heart rate is > 300 bpm | Use ECG printout to calculate heart rate. |
| | Pacing function is active | No corrective action needed. |
| Poor ECG signal quality | Poor electrode-skin contact | Reposition cable or lead wires to prevent electrodes from pulling away from patient. |
| | | Secure trunk cable clasp to patient's clothing. |
| | | Prepare skin and apply new electrodes. |
| | Outdated, corroded, or dried- out electrodes | Check Use By date on electrode packages. |
| | | Use only unexpired silver/silver chloride electrodes. Leave electrodes in sealed pouch until time of use. |
| | Loose connection. Damaged cable or | Check or reconnect cable connections. |
| | connector/lead wire | Inspect ECG and therapy cables. Replace if damaged. |
| | | Check cable with simulator and replace if malfunction observed. |
| | Noise because of radio frequency interference (RFI) | Check for equipment causing RFI (such as a radio transmitter) and relocate or turn off equipment power. |
| Baseline wander (low frequency/high amplitude artifact) | Inadequate skin preparation | Prepare skin and apply new electrodes. |
| | Poor electrode-skin contact | Check electrodes for proper adhesion. |
| | Diagnostic frequency response | Print ECG in monitor frequency response. |
| Fine baseline artifact (high frequency/low amplitude) | Inadequate skin preparation | Prepare skin and apply new electrodes. |
| | Isometric muscle tension in arms/legs | Confirm that limbs are resting on a supportive surface. |
| | | Check electrodes for proper adhesion. |

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Monitoring the ECG

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|--|--|
| ECG amplitude too small | Poor electrode-skin contact | Prepare skin and apply new electrodes. |
| | ECG lead selected | Increase ECG gain or change ECG lead. |
| | Patient condition (for example, significant myocardial muscle loss or tamponade) | Increase ECG gain or change ECG lead. |
| Monitor displays dashed lines with no ECG LEADS OFF messages | PADDLES lead selected but patient connected to ECG cable | Select one of the limb or precordial leads. |
| Monitor shows isoelectric (flat) line and PADDLES lead selected | The Test Load is connected to therapy cable | Remove the Test Load and connect therapy electrodes to cable. |
| | | Connect ECG cable and select another lead. |
| Internal pacemaker pulses difficult to see | Pacemaker pulses are very small | Turn on internal pacemaker detector (see Monitoring Patients Who Have Internal Pacemakers (on page 55). |
| | Monitor frequency response limits visibility | Connect ECG cable and select a lead other than PADDLES. |
| | | Print ECG in Diagnostic mode (see How to Print a Current Report (on page 167)). |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Acquiring a 12-Lead ECG

Intended Use

The 12-lead ECG offers paramedics and emergency physicians significant advantages over the single lead ECG trace typically available in EMS. The 12-lead ECG not only provides a diagnostic quality ECG for use in the detection of ST elevation myocardial infarction (STEMI), but also allows the knowledgeable paramedic to determine the area of myocardial injury, anticipate associated potential complications, and implement treatment strategies accordingly. In addition, the 12-lead ECG provides a baseline for serial ECG evaluations.

The 12-lead ECG transmission to the emergency department (ED) is recommended by the AHA and ERC for patients with Acute Coronary Syndrome (ACS). When transmitted from the field, 12-lead ECG has been shown to shorten time to in-hospital treatment by an estimated 10 to 60 minutes. Patients may also benefit from triage and transport to the most appropriate facility. Documentation of transient or intermittent arrhythmias and other electrophysiologic events that occur in the prehospital setting can assist in diagnosis and treatment decisions in the ED.

Indications

The 12-lead electrocardiogram is used to identify, diagnose, and treat patients with cardiac disorders and is useful in the early detection and prompt treatment of patients with acute ST-elevation myocardial infarction (STEMI).

Contraindications

None known.

12-Lead ECG Warning

WARNING

Possible Inability to Obtain a Diagnostic 12-lead ECG

Using previously unpackaged electrodes or electrodes past the Use By date may impair ECG signal quality. Remove electrodes from a sealed package immediately before use and follow the procedure for applying the electrodes.

Identifying Electrode Sites

To obtain a 12-lead ECG, place the electrodes on the limbs and the chest (precordium) as described in the following paragraph.

Limb Lead Electrode Sites

When acquiring a 12-lead ECG, limb lead electrodes are typically placed on the wrists and ankles as shown in the following figure. The limb lead electrodes can be placed anywhere along the limbs. Do not place the limb lead electrodes on the torso when acquiring a 12-lead ECG.



Figure 18 Limb Lead Electrode Placement for 12-Lead ECG

Precordial Lead Electrode Sites

The six precordial (chest) leads are placed on specific locations as shown and summarized in the following figure. Proper placement is important for accurate diagnosis and should be identified as follows: leads are V1 through V6 for AHA, or C1 through C6 for IEC. See ECG Leads Color Codes (on page 54) for color codes.

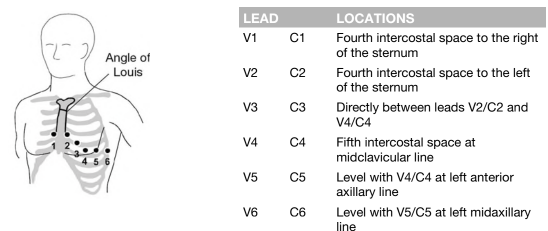


Figure 19 Precordial Lead Electrode Placement

Locating the V1/C1 position (fourth intercostal space) is critically important, because it is the reference point for locating the placement of the remaining V/C leads.

To locate the V1/C1 position:

- 1. Place your finger at the notch in the top of the sternum.
- 2. Move your finger slowly downward about 3.8 centimeters (1.5 inches) until you feel a slight horizontal ridge or elevation. This is the Angle of Louis where the manubrium joins the body of the sternum.

- 3. Locate the second intercostal space on the patient's right side, lateral to and just below the Angle of Louis.
- 4. Move your finger down two more intercostal spaces to the fourth intercostal space, which is the V1/C1 position.
- 5. Continue locating other positions from V1/C1 (see the preceding figure).

Other important considerations:

- When placing electrodes on female or obese patients, always place leads V3-V6 and C3-C6 under the breast rather than on the breast.
- Never use the nipples as reference points for locating the electrodes for men or women patients, because nipple locations vary widely.

12-Lead ECG Procedure

To acquire a 12-lead ECG:

- 1. Press ON.
- 2. Insert the lead attachments into the main cable as shown in the following figure.

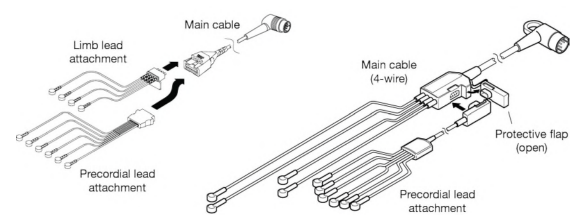


Figure 20 12-Lead ECG Cables

- 3. Insert the cable connector into the green ECG connector on the monitor.
- 4. Prepare patient's skin for electrode application (see ECG Monitoring Procedure (on page 53)).
- 5. Apply ECG electrodes (see Limb Lead Electrode Sites (on page 60)).
- 6. Encourage the patient to remain as still as possible.

WARNING

Possible Inaccurate Diagnosis

If age and sex are not entered when a 12-lead ECG is obtained, the interpretive statements are based on a default of a 50-year-old male and may provide incorrect analysis for that patient.

7. Press 12-LEAD. The 12-LEAD / AGE menu appears, prompting you to enter the patient's age.

Use the **SPEED DIAL** to select the age. Always enter the patient's age if the patient is 15 years old or younger. If you do not enter an age, the default value of 50 years is used by the interpretive analysis program and annotated on the 12-lead ECG report.

8. The 12-LEAD / SEX menu appears, prompting you to enter the patient's sex.

Use the **SPEED DIAL** to select the patient's sex. If you do not enter the sex, the default of male is used by the interpretive analysis program and is annotated on the 12-lead ECG report.

The monitor acquires, analyzes, and automatically prints the 12-lead ECG. An ECG leads-off condition for any lead is indicated on the report by a dashed line.

Note: If 15 years or less is entered for patient age, the 12-lead ECG prints at diagnostic frequency response of 0.05–150 Hz, even when 0.05–40 Hz is set up as the print default.

Note: When **12-LEAD** is pressed, internal pacemaker detection is automatically enabled, even if the function is set up to be OFF.

ECG Override

If the monitor detects signal noise while acquiring data (such as patient motion or a disconnected electrode), the screen displays the message: NOISY DATA! PRESS 12-LEAD TO ACCEPT. The message remains and 12-lead ECG acquisition is interrupted until noise is eliminated. Take appropriate action to eliminate the signal noise. This message remains as long as signal noise is detected. When signal noise is eliminated, the monitor resumes acquiring data. To override the message and acquire the 12-lead ECG in spite of the signal noise, press 12-LEAD again. The 12-lead ECG will be acquired and printed with no interpretive statements. Any 12-lead ECG report acquired in this way is annotated with the following statement: ECG OVERRIDE: DATA QUALITY PROHIBITS INTERPRETATION.

If the signal noise persists for longer than 30 seconds, 12-lead ECG acquisition stops. The screen displays **EXCESSIVE NOISE-12-LEAD CANCELLED**. You must then press **12-LEAD** to restart 12-lead ECG acquisition.

Note: If **12-LEAD** is pressed immediately after ECG electrodes are applied, the message **NOISY DATA** may occur. This message is due to the temporary instability between the electrode gel and the patient's skin that is not viewable on the ECG monitor screen, but is detected as noisy data. In general, it is best to wait at least 30 seconds after applying the last electrode before pressing the **12-LEAD** button, to allow for electrode/skin stabilization. Also, good skin preparation shortens the stabilization time.

Computerized ECG Analysis

Computerized ECG analysis statements are automatically printed on 12-lead ECG reports. Printing of the interpretive statements is a setup option and may be turned off in Setup mode. For information on how to change this setup option, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

Chapter 4 | Monitoring

The interpretative statements pertaining to myocardial injury, infarct, and ischemia are derived from measurements made on a signal-averaged beat (median beat) formed for each of the 12 leads. The computerized ECG analysis selects three representative beats from the ten seconds of data for each lead and averages the three beats to derive the median beat for that lead. The ECG analysis is always based on ECG data obtained at 0.05–150 Hz frequency response.

The analysis program is adjusted for patient age and sex. The 12-lead ECG interpretive algorithm used by the LIFEPAK 15 monitor/defibrillator is the University of Glasgow 12-Lead ECG Analysis Program. For more information, contact your Physio-Control representative for a copy of the *Physio-Control Glasgow 12-Lead ECG Analysis Program Physician's Guide*.

WARNING

Possible Incorrect Treatment with Reperfusion Therapy

Computerized ECG interpretive statements should not be used to withhold or prescribe patient treatment without review of the ECG data by qualified medical personnel. All 12-lead ECG interpretation statements provided by the LIFEPAK 15 monitor/defibrillator include the printed message **UNCONFIRMED**. Always confirm interpretive statements by over-reading the ECG data.

Printed 12-Lead ECG Report Formats

Two 12-lead ECG report formats are available for printing: 3-channel or 4-channel. In addition, each of those formats can be printed in standard and cabrera styles.

The 3-Channel Format

The 3-channel format prints 2.5 seconds of data for each lead. The following figure is an example of a 12-lead ECG report printed in the 3-channel format, standard style. The following figure is an example of a 12-lead ECG report printed in the 3-channel format, cabrera style. The sequence in which the limb leads are presented differs between the standard and cabrera styles, as shown. The default format for printing 12-lead ECG reports is 3-channel standard. To change the printed format of 12-lead ECG reports, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device. Alternatively, press **OPTIONS**, select **PRINT**, select **REPORT**: 12-LEAD, and then select **FORMAT**.

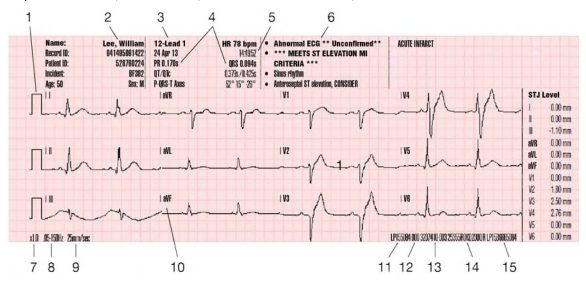


Figure 21 Example of Printed 3-Channel, Standard 12-Lead ECG Report

| FIG | URE LEGEND | | |
|-----|----------------------------|----|--------------------|
| 1 | 1 mV reference | 9 | Printer speed |
| 2 | Patient ID | 10 | Lead annotation |
| 3 | Report type and number | 11 | Device number |
| 4 | Standard measurement | 12 | Site number |
| 5 | Time/date 12-lead acquired | 13 | Software version |
| 6 | Computerized ECG analysis | 14 | Configuration code |
| 7 | ECG size | 15 | Serial number |
| 8 | Frequency response | | |

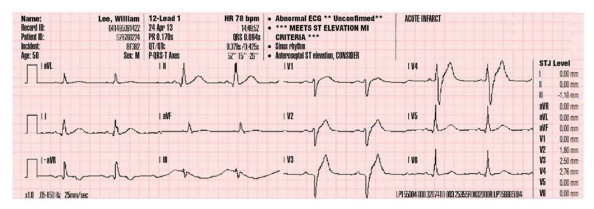


Figure 22 Example of Printed 3-Channel, Cabrera 12-Lead ECG Report

The 4-Channel Format

The following two figures are examples of 12-lead ECG reports printed in the 4-channel format. The 4-channel format consists of the median complex (or median beat) derived for each of the 12 leads and 10 seconds of data for Lead II.

Note: The fiducial marks displayed in the 4-channel format identify the measurement intervals used for the interpretive statements of the analysis program. These marks are part of the analysis program and cannot be turned off.

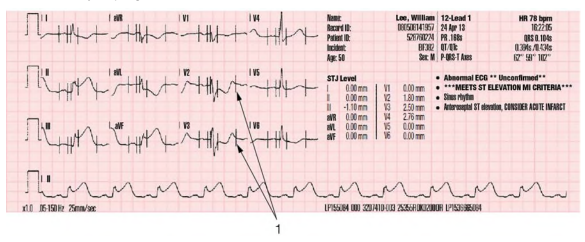


Figure 23 Example of Printed 4-Channel, Standard 12-Lead ECG Report

FIGURE LEGEND

1 Fiducial marks

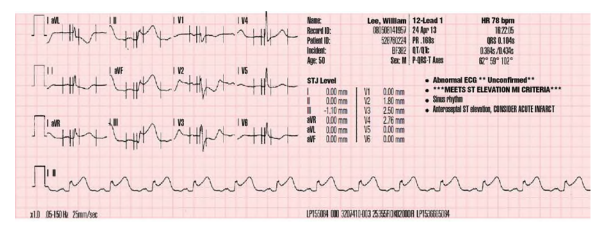


Figure 24 Example of Printed 4-Channel, Cabrera 12-Lead ECG Report

Printed 12-Lead ECG Frequency Response

The 12-lead ECG can be printed in two diagnostic frequency responses (or bandwidths): 0.05–40 Hz and 0.05–150 Hz. The frequency response of 0.05–150 Hz is the Association for the Advancement of Medical Instrumentation (AAMI) standard for diagnostic ECGs. The 0.05–40 Hz setting preserves the low frequency limit that is needed for the diagnosis of myocardial ischemia and infarction while reducing high frequency artifact (in particular from patient muscle tension) to help make the diagnostic printout less noisy and more readable.

Note: The LIFEPAK 15 monitor/defibrillator acquires ECG data and performs the interpretive analysis based on the full frequency of 0.05–150 Hz. The 0.05–40 Hz bandwidth affects only the printed appearance of the ECG data.

The 12-lead ECG printed in the 0.05–40 Hz setting can be used to diagnose acute myocardial ischemia and ST-segment elevation myocardial infarction (STEMI). This is because the low frequency limit of 0.05 Hz is not changed from the standard diagnostic setting of 0.05–150 Hz. The 0.05 Hz frequency provides accurate representation of low frequency signals, that is, the P, ST segment, and T waves. The presence or absence of ST segment changes indicative of myocardial ischemia or infarction will be accurately reproduced. In addition, the criteria for visual analysis and interpretation of cardiac rhythm and PR, QRS, and QT intervals are preserved, as is true with hospital cardiac monitors that have an upper frequency limit of 40 Hz.

However, in some adult patients, the amplitude (that is, voltage) of the QRS may be reduced when 12-lead ECGs are printed at the upper limit of 40 Hz rather than at 150 Hz. Therefore, certain diagnoses, which depend on R wave amplitude (for example, ventricular hypertrophy), should not be made using this setting. In the pediatric patient, this effect on R wave amplitude is particularly noticeable because QRS durations in children are typically quite narrow. Because R wave amplitude reduction is more likely with pediatric patients, the 12-lead ECG automatically prints at 0.05–150 Hz, overriding the 40 Hz limit, when a patient age of 15 years or younger is entered.

Troubleshooting Tips

Table 12 Troubleshooting Tips for the 12-Lead ECG

| Table 12 Troubleshooti | ng Tips for the 12-Lead ECG | |
|--|--|---|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
| Any of these messages displayed: | One or more ECG electrodes disconnected | Confirm ECG electrode connections. |
| CONNECT ECG LEADS | ECG cable is not connected to monitor | Confirm ECG cable connections. |
| ECG LEADS OFF XX LEADS OFF | Poor electrode-skin contact | Reposition cable and/or lead wires to prevent electrodes from pulling away from patient. |
| | | Secure trunk cable clasp to patient's clothing. |
| | | Prepare skin and apply new electrodes. |
| | Broken lead wire | Select another lead. |
| | | Select PADDLES lead, and use standard paddles or therapy electrodes for ECG monitoring. |
| | | Check ECG cable continuity. |
| Noisy signal and/or message displayed: NOISY DATA! PRESS 12-LEAD TO ACCEPT | Noise in a lead other than the displayed lead | Press 12-LEAD again to override the message. Examine the printout to determine leads affected by noise. Replace or reposition the affected electrodes and lead wires. |
| | Poor electrode-skin contact | Reposition cable and/or lead wires to prevent electrodes from pulling away from patient. |
| | | Secure trunk cable clasp to patient's clothing. |
| | | Prepare skin and apply new electrodes. |
| | Loose connection | Check or reconnect cable connections. |
| | Patient motion | Encourage patient to lie quietly.Support patient's limbs. |
| | Vehicle motion | Stop vehicle while acquiring 12-lead ECG data. |
| | Outdated, corroded, or dried-out electrodes | Check Use By date on electrode packages. Use only unexpired silver/silver chloride electrodes. Leave electrodes in sealed pouch until time of use. |
| | Radio Frequency Interference (RFI) | Check for equipment causing RFI (such as a radio transmitter) and relocate or turn off equipment power. |
| | Damaged cable or connector/lead wire | Inspect main cable and attachments. Replace if damaged. |
| Monitor does not complete 12-lead ECG operation sequence | Operator pressed another function button (such as PRINT) before 12-lead ECG sequence completed | Press 12-LEAD to acquire another 12-lead ECG. Allow enough time for sequence to complete. |

Monitoring SpO2, SpCO, and SpMet

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|---------------------------------------|--|
| Noisy signal and message displayed: EXCESSIVE NOISE- 12-LEAD CANCELLED | Signal noise for more than 30 seconds | Press 12-LEAD to acquire another 12-lead ECG. |
| Baseline wander (low frequency/high amplitude artifact) | Inadequate skin preparation | Prepare skin as described in ECG Monitoring Procedure (on page 53) and apply new electrodes. |
| | Poor electrode-skin contact | Check electrodes for proper adhesion. |
| Fine baseline artifact (high frequency/low amplitude) | Inadequate skin preparation | Prepare skin as described in ECG Monitoring Procedure (on page 53) and apply new electrodes. |
| | Isometric muscle tension in arms/legs | Confirm that limbs are resting on a supportive surface. |
| | | Check electrodes for proper adhesion. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Monitoring SpO2, SpCO, and SpMet

SpO₂, SpCO™, and SpMet™ are optional features for the LIFEPAK 15 monitor/defibrillator. When all three options (SpO₂, SpCO, and SpMet) are installed, the pulse oximeter measures functional oxygen saturation (SpO₂), carboxyhemoglobin concentration (SpCO), and methemoglobin concentration (SpMet) in the blood.

IMPORTANT! SpO₂-only sensors and combination SpO₂, SpCO, and SpMet sensors are available for use. Masimo[®] SpO₂-only sensors that have a red connector are compatible with the LIFEPAK 15 monitor. Masimo Rainbow[®] sensors are necessary to monitor SpCO and SpMet in addition to SpO₂. These sensors are not compatible with other LIFEPAK defibrillator/monitors.

Nellcor SpO₂ sensors may be used with the LIFEPAK 15 monitor/defibrillator, if the Masimo Red™ MNC adapter cable is used.

For a list of SpO₂ sensors and connector cables that are intended for use with the LIFEPAK 15 monitor/defibrillator, see the Physio-Control website. Carefully read the Directions for Use that are provided with the sensors and connector cables for a complete description, instructions, warnings, cautions, and specifications. To order sensors and connector cables, contact your Physio-Control representative. In the USA, call Customer Support at 1.800.442.1142, option 2.

Intended Use

A pulse oximeter is a noninvasive device that continuously measures functional oxygen saturations (SpO₂), carboxyhemoglobin concentration (SpCO), and methemoglobin concentration (SpMet) in the blood. Continuously monitoring SpO₂ can provide an early warning when oxygen saturation is decreasing and can help the clinician act rapidly before the patient develops the later signs of hypoxemia. Previously, the blood parameters SpCO and SpMet could only be obtained from invasive blood gas samples. This new technology assists in identifying the

often hidden conditions of carboxyhemoglobinemia (carbon monoxide poisoning) and methemoglobinemia (a condition that impedes delivery of oxygen to the tissues). Low levels of both SpCO and SpMet are normally found in the blood; however, early detection of significantly high levels can lead to proper diagnosis and treatment, and can help improve patient outcome.

Pulse oximetry is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times; do not rely solely on the SpO₂, SpCO, and SpMet measurements. If a trend toward patient deoxygenation is evident or carbon monoxide poisoning or methemoglobinemia is suspected, blood samples should also be analyzed using laboratory instruments to completely understand the patient's condition.

Do not use the pulse oximeter to monitor patients for apnea, or as a replacement or substitute for ECG-based arrhythmia analysis.

Indications

Pulse oximetry is indicated for use in any patient who is at risk of developing hypoxemia, carboxyhemoglobinemia, or methemoglobinemia. SpO₂ monitoring may be used during no motion and motion conditions, and in patients who are well or poorly perfused. SpCO and SpMet accuracies have not been validated under motion or low perfusion conditions.

Contraindications

None known.

SpO2, SpCO, and SpMet Warnings and Cautions

| WARNING | Shock or Burn Hazard |
|---------|--|
| | Before use, carefully read these operating instructions the sensor and cable directions for use, and precautionary information. |
| WARNING | Shock or Burn Hazard |
| | Using other manufacturers' sensors or cables may cause improper oximeter performance and invalidate safety agency certifications. Use only sensors and cables that are specified in these operating instruction |
| WARNING | Inaccurate Pulse Oximeter Readings |
| | Do not use a damaged sensor or cable. Do not alter the sensor or cable in any way. Alterations or modification may affect performance and/or accuracy. Never use more than one cable between the pulse oximeter and the sensor to extend the length. |

WARNING

Inaccurate Pulse Oximeter Readings

Sensors exposed to ambient light when incorrectly applied to a patient may exhibit inaccurate saturation readings. Securely place the sensor on the patient and check the sensor's application frequently to help ensure accurate readings.

WARNING

Inaccurate Pulse Oximeter Readings

Severe anemia, hypothermia, severe vasoconstriction, carboxyhemoglobin, methemoglobin, intravascular dyes that change usual blood pigmentation, elevated bilirubin, excessive patient movement, venous pulsations, electrosurgical interference, exposure to irradiation and placement of the sensor on an extremity that has a blood pressure cuff, intravascular line, or externally applied coloring (such as nail polish) may interfere with oximeter performance. The operator should be thoroughly familiar with the operation of the oximeter prior to use.

WARNING

Inaccurate Pulse Oximeter Readings

The pulsations from intra-aortic balloon support can be additive to the pulse rate on the oximeter pulse rate display. Verify patient's pulse rate against the ECG heart rate.

WARNING

Possible Skin Injury

Prolonged, continuous use of a sensor may cause irritation, blistering, or pressure necrosis of the skin. Check the sensor site regularly based on patient condition and type of sensor. Change the sensor site if skin changes occur. Do not use tape to hold the sensor in place as this may cause inaccurate readings or damage to the sensor or skin.

WARNING

Possible Strangulation

Carefully route patient cabling to reduce the possibility of patient entanglement or strangulation.

CAUTION

Possible Equipment Damage

To avoid damage to the cable, always hold by the connector rather than the cable, when connecting or disconnecting either end.

CAUTION

Possible Equipment Damage

Do not soak or immerse the sensors or cables in any liquid solution. Do not attempt to sterilize.

No Implied License

Possession or purchase of the pulse oximeter does not convey any expressed or implied license to use the pulse oximeter with unauthorized sensors or cables which would, alone or in combination with this device, fall within the scope of one or more of the patents relating to this device.

How a Pulse Oximeter Works

A pulse oximeter sensor directs light through a patient's fleshy body site (usually a finger or toe). The sensor sends wavelengths of light from the emitter to the receiving detector as shown in the following figure.

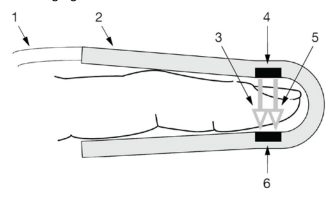


Figure 25 How a Pulse Oximeter Works

FIGURE LEGEND

- 1 Cable
- 2 Sensor (holds LEDs and detector)
- 3 Red

- 4 Light-emitting diodes
- 5 Infrared
- 6 Light-receiving detector

The pulse oximeter translates the amount of light received by the detector to the various forms of hemoglobin saturation levels and displays them as SpO₂, SpCO, and SpMet percentages. Normal values for SpO₂ typically range from 95% to 100%. Normal values for SpCO are typically less than 9% (the higher range of normal is often seen in smokers). Normal values for SpMet are typically less than 2% and may be caused by exposure to some pharmaceuticals including local anesthetic agents and chemical agents such as nitrites.

SpO2, SpCO, and SpMet Monitoring Considerations

The quality of the SpO₂, SpCO, and SpMet readings depends on correct sensor size and placement, adequate blood flow through the sensor site, and limiting patient motion and sensor exposure to ambient light. For example, with very low perfusion at the sensor site, readings may be lower than core arterial oxygen saturation. Test methods for accuracy are available by contacting your local Physio-Control representative.

Use the following criteria to select the appropriate pulse oximeter sensor:

- · Patient size (adult, pediatric, infant) and weight
- Patient perfusion to extremities
- Patient activity level
- Available application sites on the patient's body
- Sterility requirements
- Anticipated duration of monitoring

To help ensure optimal performance:

- Use a dry and appropriately sized sensor.
- Choose a site that is well perfused. The ring finger is preferred.
- Choose a site that least restricts patient movement, such as finger of the non-dominant hand.
- Be sure the fleshy part of the digit completely covers the detector.
- Keep the sensor site at the same level as the patient's heart.
- Apply the sensor according to the Directions for Use provided with the sensor.
- Observe all warnings and cautions noted in the sensor's Directions for Use.

Sensor Application

The preferred site for sensor application is the ring finger of the non-dominant hand. To position the sensor:

- 1. Orient the sensor so the cable is on the back of the patient's hand.
- 2. Place the finger in the sensor until the tip of the finger touches the "raised digit stop."
- The hinged tabs of the sensor should open to evenly distribute the grip pressure of the sensor along the length of the finger. Check the arrangement of the sensor to verify correct positioning. Complete coverage of the detector window is needed to ensure accurate data.

The sensors are sensitive to light. If excessive ambient light is present, remove or reduce lighting, cover the sensor site with an opaque material to block the light, and check appropriateness of sensor site. Failure to do so could result in inaccurate measurements.

If excessive movement presents a problem during SpCO/SpMet monitoring, consider the following possible solutions:

- Be sure the sensor is secure and properly aligned.
- Use a disposable adhesive sensor.
- If possible, move the sensor to a less active site.

Note: Wrapping the sensor too tightly or using supplemental tape to hold the sensor in place may cause inaccurate oximeter readings.

Note: Circulation distal to the sensor site should be checked routinely.

IMPORTANT! Masimo Rainbow sensors are necessary to monitor SpCO and SpMet and are not compatible with other LIFEPAK defibrillator/monitors.

Oximeter Monitoring Procedure

Power to the pulse oximeter is controlled by the LIFEPAK 15 monitor/defibrillator. When the defibrillator is turned on, the oximeter turns on and performs a calibration and self-test that requires approximately 20 seconds. During the calibration and self-test, the screen does not display SpO₂, SpCO, or SpMet information.

To conserve battery power, the pulse oximeter goes into "sleep mode" when not in use. Sleep mode is activated within 10 seconds of disconnecting the sensor. During sleep mode, the screen does not display SpO₂, SpCO, or SpMet information. When a sensor or patient signal is detected, the oximeter performs a self-test and then returns to normal mode.

The pulse oximeter measures and displays SpO_2 levels between 50 and 100%. SpO_2 levels less than 50% are displayed as <50. The pulse oximeter measures and displays SpCO in the range of 0–40%. The pulse oximeter measures and displays SpMet in the range of 0–15%. Measurement accuracies are specified in the $SpO_2/SpCO/SpMet$ section in Appendix A.

To monitor SpO₂:

- 1. Press ON.
- 2. Connect the pulse oximeter cable to the monitor and sensor.
- 3. Attach the sensor to the patient.
- 4. Observe the pulse bar for fluctuation. Amplitude of the pulse bar indicates relative signal quality.
- 5. Confirm that the SpO₂ reading appears and is stable.
- 6. Use the SPEED DIAL to adjust volume, sensitivity, and averaging time, as necessary.

Monitoring SpO2, SpCO, and SpMet

To monitor SpCO or SpMet:

- 1. Perform Step 2 through Step 5 above.
- 2. Verify that an SpCO/SpMet sensor is in use. Only Rainbow sensors are capable of reading SpCO/SpMet.
- Encourage the patient to remain still.
- To quickly obtain SpCO or SpMet value, press PRINT. If dashes (---) appear on printout instead of values for SpCO or SpMet, allow a few more seconds for measurement to be obtained.

or

To display SpCO or SpMet:

- Use the SPEED DIAL to select the SpO2 area.
- Select PARAMETER from menu.
- Select **SPCO** or **SPMET**. Selected value displays for 10 seconds.

Note: SpCO and SpMet monitoring are not intended for use under patient motion or low perfusion conditions.

SpCO/SpMet Advisory

If the SpCO or SpMet reading is above normal limits, indicating a dangerous amount of carboxyhemoglobin or methemoglobin, an Advisory occurs.

During an Advisory:

- The elevated SpCO or SpMet value is displayed instead of SpO₂.
- The elevated value flashes and the alarm tone sounds.
- One of the following Advisory messages appears in the message area:

Advisory: SpCO > 10%

Advisory: SpMet > 3%

To cancel the Advisory, press **ALARMS**. The SpO₂ area reverts to the SpO₂ reading. The Advisory message remains on the screen until the elevated value returns to within normal limits or the device is turned off.

WARNING

Inaccurate SpO₂ Readings

Carboxyhemoglobin and methemoglobin may erroneously increase SpO₂ readings. The amount that SpO₂ increases is approximately equal to the amount of carboxyhemoglobin or methemoglobin that is present.

WARNING

Inaccurate SpCO and SpMet Readings

Very low arterial oxygen saturation levels may cause inaccurate SpCO and SpMet readings.

The Pleth Waveform

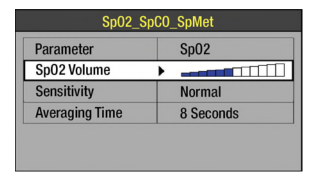
You can display the plethysmographic (pleth) waveform in Channel 2 or 3.

To display the pleth waveform:

- 1. Rotate the SPEED DIAL to outline waveform CHANNEL 2 or 3.
- 2. Press the **SPEED DIAL**. The Channel menu appears.
- 3. Select **WAVEFORM** and then select **SPO2**. The SpO₂ waveform appears in the selected channel. The waveform is automatically sized for optimum waveform viewing.

Volume

To adjust the pulse tone volume:



- 1. Rotate the **SPEED DIAL** to outline the SpO₂ area on the Home Screen.
- 2. Press the SPEED DIAL.
- 3. Highlight and select SPO2 VOLUME.
- 4. Rotate the **SPEED DIAL** to the desired volume.
- Press the SPEED DIAL to set the volume.

Sensitivity

The sensitivity setting allows you to adjust the oximeter to either **NORMAL** or **HIGH** for differing perfusion states.

To adjust sensitivity:

- 1. Outline and select the SpO₂ area on the Home Screen.
- 2. Select **SENSITIVITY** and then select **NORMAL** or **HIGH**.

Note: NORMAL sensitivity is recommended for most patients. The **HIGH** sensitivity setting allows SpO₂ monitoring under low perfusion states, such as the severe hypotension of shock. However, when SpO₂ sensitivity is set to **HIGH**, the signal is more susceptible to artifact. Monitor the patient closely when using the **HIGH** sensitivity setting.

Averaging Time

Averaging time allows you to adjust the time period that is used to average the SpO₂ value.

To adjust averaging time:

- 1. Outline and select the SpO₂ area on the Home Screen.
- 2. Select AVERAGING TIME and then select one of the following:
 - 4 Seconds
 - 8 Seconds
 - 12 Seconds
 - 16 Seconds

Note: Averaging time of 8 seconds is recommended for most patients. For patients with rapidly changing SpO₂ values, 4 seconds is recommended. Use a 12- or 16-second time period when artifact is affecting the performance of the pulse oximeter.

Pulse Rate Monitoring

If ECG monitoring is not active, the SpO₂ sensor can be used to monitor the patient's pulse rate. The pulse rate value is indicated by **PR** (**SPO2**).

Pulse rate monitoring is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times. Check pulse manually if patient shows signs of abnormal pulse rate.

Cleaning

Pulse oximetry sensors may be adhesive (single-patient use) or reusable.

To clean the reusable sensor and connector cable:

- 1. Disconnect the sensor and cable from the monitor. Inspect the cable for damage.
- 2. Use a clean, soft cloth dampened with 70% isopropyl alcohol to wipe clean.
- 3. Allow to dry thoroughly before placing the sensor on a patient or reconnecting the cable to the monitor.

Note: Do not attempt to sterilize. Do not soak or immerse in any liquid solution. For information about cleaning the device, see Cleaning the Device (on page 212).

Troubleshooting Tips

Table 13 Troubleshooting Tips for SpO2, SpCO, and SpMet

| POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|--|
| Excessive patient motion | Keep patient still.Check that sensor is secure.Relocate sensor.Apply adhesive sensor. |
| Patient perfusion may be too low | Check patient.Increase sensitivity. |
| Excessive patient motion | Keep patient still. Check that sensor is secure. Relocate sensor. Apply adhesive sensor. Increase sensitivity. |
| An electrosurgical unit (ESU) may be interfering with performance | Move the monitor as far as possible from the ESU. Plug the ESU and monitor into different circuits. Move the ESU ground pad as close to the surgical site as possible. |
| Sensor may be damp | Replace sensor. |
| Sensor not connected to patient or cable disconnected from monitor/defibrillator | Check that sensor and cable are connected properly. Check that appropriate sensor is in use. |
| Damaged cable or sensor | Replace damaged cable or sensor. |
| Sensor may be too tight | Reposition sensor.Relocate sensor. |
| Patient is in cardiac arrest or shock | Check patient. |
| Oximeter may be performing self-calibration or self-test | Wait for completion. If values do not display within 30 seconds, disconnect and reconnect sensor. If values do not display within another 30 seconds, replace sensor. |
| Defibrillator shock just delivered | None. If values do not display within 30 seconds, disconnect and reconnect sensor. If values do not |
| | Patient perfusion may be too low Excessive patient motion An electrosurgical unit (ESU) may be interfering with performance Sensor may be damp Sensor not connected to patient or cable disconnected from monitor/defibrillator Damaged cable or sensor Sensor may be too tight Patient is in cardiac arrest or shock Oximeter may be performing self-calibration or self-test |

Monitoring SpO2, SpCO, and SpMet

| OBSERVATION | DOSSIDI E CAUSE | CODDECTIVE ACTION |
|--|--|---|
| OBSERVATION | POSSIBLE CAUSE High intensity lights (such as | CORRECTIVE ACTION |
| | pulsating strobe lights) may be interfering with performance | Cover sensor with opaque material, if necessary. |
| | Damaged cable or sensor | Replace damaged cable or sensor. |
| Different SpCO or SpMet measurements on same patient | Every measurement, even on the same patient, can be different | Confirm by taking three measurements: ring finger, middle finger, and then index finger; average the results. |
| XXX appears in place of SpO₂ reading | SpO₂ module failed. Internal cable failed. | Turn device off and then on again. If problem persists, contact qualified service personnel. |
| SPO2: CHECK SENSOR message appears | Sensor is disconnected from patient or cable | Attach the sensor.Check that sensor is secure. |
| | Excessive ambient light | Remove or block light source, if possible. |
| | | Cover sensor with opaque material, if necessary. |
| | Faulty or defective sensor | Replace sensor. |
| | Patient has a weak pulse or low blood pressure, or the sensor is not properly placed | Change sensor location.Check if patient perfusion is adequate for sensor location. |
| | | Check that sensor is secure and not too tight. |
| | | Check that sensor is not on extremity with blood pressure cuff or intravascular line. |
| | | Test sensor on someone else. |
| SPO2: UNKNOWN SENSOR message appears | A sensor that is not Physio-Control approved is | Check that the sensor is approved by Physio-Control. |
| | connected to the device | If using Nellcor sensor, check that it is connected to monitor using Masimo Red MNC adapter cable. |
| SPO2: SEARCHING FOR PULSE message appears | A sensor is connected to the patient and is searching for a pulse | Wait for completion. |
| SPO2: LOW PERFUSION message appears | Patient has a weak pulse | Change sensor location. |
| SP02: POOR QUALITY SIGNAL message appears | When the signal quality is low, the accuracy of the measurement may be compromised | Check that sensor and cable are connected properly. Move sensor to a better perfused site. |
| SPCO: POOR QUALITY SIGNAL message appears | When the signal quality is low, the accuracy of the measurement may be compromised | Check that sensor and cable are connected properly. Move sensor to a better perfused site. |
| | | |

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|--|---|
| SPMET: POOR QUALITY SIGNAL message appears | When the signal quality is low, the accuracy of the measurement may be compromised | Check that sensor and cable are connected properly. Move sensor to a better perfused site. |
| SPCO/SPMET: POOR QUALITY SIGNAL message appears | When the signal quality is low, the accuracy of the measurement may be compromised | Check that sensor and cable are connected properly. Move sensor to a better perfused site. |
| SPO2: SENSOR DOES NOT SUPPORT SPCO OR SPMET message appears | SpO ₂ -only sensor used with SpCO/SpMet capable device | None necessary, or use Rainbow sensor to measure SpCO or SpMet. |

Note: Most Rainbow sensor messages (SpO₂, SpCO, and SpMet) are reported as **SPO2**: **(MESSAGE)**. The **POOR QUALITY SIGNAL** message indicates the specific parameter affected.

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Monitoring Noninvasive Blood Pressure

Intended Use

The LIFEPAK 15 noninvasive blood pressure (NIBP) monitor measures blood pressure (BP) using the oscillometric measurement technique to determine systolic, diastolic, and mean arterial pressures, and pulse rate. The measurement can be initiated manually or set to recur automatically at predetermined intervals.

Blood pressure measurements determined using this device are equivalent to those obtained by a trained observer using the cuff/stethoscope auscultation method, within the limits prescribed by the American National Standard *Electronic or automated sphygmomanometers* (AAMI SP-10).

NIBP is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times; do not rely solely on the NIBP monitor.

Indications

Noninvasive blood pressure monitoring is intended for detection of hypertension or hypotension and monitoring BP trends in patient conditions such as, but not limited to, shock, acute dysrhythmia, or major fluid imbalance. NIBP monitoring is not indicated for neonatal patients less than one month old.

Contraindications

None known.

NIBP Monitoring Warnings and Caution

| WARNING | Possible Loss of Intravenous Access and Inaccurate Infusion Rate |
|---------|---|
| | Do not apply the blood pressure cuff on an extremity that is used for an intravenous infusion or arteriovenous (A-V) shunt. Patency of the intravenous infusion may be affected by blood pressure measurement due to the occlusion of blood flow. |
| WARNING | Possible Circulation Impairment |
| | Blood flow to the extremity may be impaired by prolonged, continuous use of a blood pressure cuff, a kink in the tubing, or frequent measurements. Check circulation regularly and loosen or reposition the cuff if changes in circulation occur. |
| WARNING | Possible Inaccurate Blood Pressure Readings |
| | Do not alter the NIBP monitor's pneumatic tubing. Altering NIBP tubing may cause improper performance and may void the warranty. Avoid compression or restriction of pressure tubes. |
| WARNING | Possible Patient Harm |
| | Do not apply the blood pressure cuff over a wound. Doing so may cause further injury. |
| WARNING | Possible Patient Harm |
| | Do not apply the blood pressure cuff on the arm on the side of a mastectomy. |
| WARNING | Possible Inaccurate Blood Pressure Readings |
| | Using NIBP accessories not recommended by Physio-Control may cause the device to perform improperly and invalidate the safety agency certifications. Use only the accessories that are specified in these operating instructions. |
| WARNING | Possible Inaccurate Oxygen Saturation Readings |
| | Do not perform NIBP measurement on an extremity used for oxygen saturation monitoring. Oxygen saturation measurement is affected by blood pressure |

measurement due to the occlusion of blood flow.

CAUTION

Equipment Damage

Do not inflate a cuff unless it is placed on an extremity.

How NIBP Monitoring Works

The NIBP monitor uses the oscillometric measurement technique. The oscillometric technique does not use Korotkoff sounds to determine blood pressure; rather, it monitors the changes in pressure pulses that are caused by the flow of blood through the artery. The NIBP monitor inflates the cuff around the patient's arm to a value that occludes the artery, and then deflates the cuff in steps. When blood starts to flow through the artery, the increasing blood flow causes the amplitude of the pressure pulses in the cuff to increase. As the NIBP monitor steps the pressure down, the pulses reach a peak amplitude and then start to decrease. The rising and falling amplitude values form a curve that is analyzed to yield systolic pressure, diastolic pressure, and mean arterial pressure (MAP).

The NIBP monitor measures the pulse rate by tracking the number of pulses over time. The NIBP monitor uses artifact rejection techniques to provide accurate results under most operating conditions. When a patient is experiencing arrhythmias during a measurement, the accuracy of the pulse determination may be affected or the time needed to complete a measurement may be extended. In shock conditions, the low amplitude of blood pressure waveforms makes it difficult for the monitor to accurately determine the systolic and diastolic pressures.

NIBP Monitoring Considerations

As with any noninvasive oscillometric blood pressure monitor, clinical conditions can affect the accuracy of the measurements obtained, including the following:

- The patient's physiological condition. For example, shock may result in a blood pressure
 waveform that has a low amplitude, making it difficult for the monitor to accurately determine
 the systolic and diastolic pressures. Altered hemodynamics caused by pregnancy, including
 preeclampsia, may result in inaccurate readings.
- The position of the patient.
- Motion may prolong the measurement process since motion artifacts have to be rejected in the data stream. Motion that affects measurement can include patient movement, patient seizure, bumping the cuff, and flexing the extremity under the cuff.
- The presence of other medical devices. The NIBP monitor does not operate effectively if the patient is connected to a heart/lung machine.
- Extremes of temperature, humidity, or altitude.
- When a patient is experiencing arrhythmias, pulse rate accuracy may be affected or the time needed to complete an NIBP measurement may be extended. The device automatically deflates if a blood pressure measurement cannot be obtained in 120 seconds.
- Blood pressure and pulse can fluctuate greatly between measurements; the monitor cannot alert the operator of changes in vital signs that occur between measurement cycles.

Monitoring Noninvasive Blood Pressure

- There may be some difference between readings taken manually and readings from the NIBP monitor due to the differing sensitivity of the two methods. The NIBP monitor meets the ANSI/SP10 AAMI standard that requires a mean difference of ±5 mmHg, with a standard deviation no greater than 8 mmHg, compared to auscultatory readings.
- When using the NIBP monitor during defibrillation, the NIBP monitor is not available when the defibrillator is being charged. Upon shock, the monitor resets and dashes (- -) appear in place of pressure readings. After defibrillation, you can resume blood pressure measurement according to NIBP Monitoring Procedure (on page 82).
- If the blood pressure cuff fails to deflate for any reason or causes undue discomfort to the patient, remove the cuff from the arm or disconnect the tubing from the defibrillator.
- If the patient has been active, optimal resting measurements will be obtained if you wait five minutes before taking a blood pressure measurement.

Cuff Selection

The use of properly designed and sized cuffs is essential for the accurate measurement of blood pressure. The cuff must fit snugly around the extremity to occlude the artery. For a list of BP cuffs that are intended for use with the LIFEPAK 15 monitor/defibrillator, see the LIFEPAK 15 Monitor/Defibrillator Accessories Catalog at www.physio-control.com.

NIBP Monitoring Procedure

The NIBP monitor inflates an occluding cuff and determines systolic and diastolic pressures, mean arterial pressure (MAP), and pulse rate. Pressure measurements are reported in mmHg and pulse rate in beats per minute (bpm).

Both single-measurement and specified-interval (timer-controlled) methods of blood pressure reading are available.

The NIBP monitor draws power from the defibrillator. When the defibrillator is turned on, the NIBP monitor conducts a self-test that takes approximately three seconds.

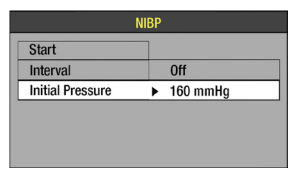
IMPORTANT! The LIFEPAK 15 monitor NIBP port and tubing are not compatible or interchangeable with the NIBP tubing that is used with other LIFEPAK monitor/defibrillators.

Changing the Initial Inflation Pressure

The initial cuff pressure should be set approximately 30 mmHg higher than the patient's anticipated systolic pressure. The factory default initial inflation pressure for the first measurement is 160 mmHg. For pediatric patients, the initial cuff pressure may need to be lowered. Initial inflation settings are 80, 100, 120, 140, 160, or 180 mmHg. For infants, the recommended initial cuff pressure is 120 mmHg.

Caution should be taken not to lower the initial pressure below the adult patient's systolic measurement. Doing so may cause the cuff to reinflate and cause patient discomfort. For subsequent measurements, the monitor inflates approximately 30 mmHg higher than the previously determined systolic pressure.

To select an initial pressure:



- Rotate the **SPEED DIAL** to outline the NIBP area
- 2. Press the **SPEED DIAL**. The NIBP menu appears.
- 3. Select INITIAL PRESSURE.
- 4. Rotate the **SPEED DIAL** to the desired pressure.
- 5. Press the **SPEED DIAL** to set the initial pressure.

Note: Measurement data is recorded in the LIFEPAK 15 monitor/defibrillator Vital Sign Log. For more information about the Vital Sign Log and its use, see Data Management (on page 159).

Manual Single-Measurement Procedure

The NIBP measurement typically takes 40 seconds to complete. If the measurement is not completed within 120 seconds, the cuff automatically deflates.

To obtain a manual single measurement:

- 1. Press ON.
- 2. Select the appropriately-sized cuff.
- 3. Properly align the cuff artery markings, if present, and apply snugly to the extremity.
- 4. Connect the tubing to the cuff and to the NIBP port on the monitor.
- 5. Change the initial inflation pressure, if necessary.
- 6. If possible, ensure the patient is comfortably seated with feet flat on the floor, legs uncrossed, and back supported. Ask the patient to relax as much as possible and refrain from talking during the measurement procedure. The operator should be able to view the device screen during the measurement.
- 7. Position the extremity in a relaxed and supported position at approximately the same level as the right atrium of the patient's heart. Inform the patient that the cuff will inflate and cause a "big squeeze" around the arm and that the patient's fingers may tingle.
- Press NIBP to start the measurement, and check that the patient's arm is not moving.
 When the measurement is complete, systolic, diastolic, and mean arterial pressures are displayed.

To cancel a measurement, press **NIBP** again.

Note: NIBP pulse rate is displayed only when ECG or SpO₂ is not active.

Timer-Controlled Measurement Procedure

When the timer is set, the monitor performs recurring measurements at a fixed interval. When using timer-controlled measurement, the interval is counted from the start of the measurement to the start of the next measurement. Choices are **OFF** (factory default), **2**, **3**, **5**, **10**, **15**, **30**, and **60** minutes.

To take a manual measurement between timer-controlled measurements, press **NIBP**. The next interval is counted from the beginning of the manual measurement.



Figure 26 NIBP Measurements and Timer

FIGURE LEGEND

- 1 Countdown timer-displays time until next measurement
- 2 Mean arterial pressure (MAP)
- 3 Systolic pressure
- 4 Diastolic pressure

To set timer-controlled measurements:

- 1. Press ON.
- Select the appropriately-sized cuff.
- 3. Properly align the cuff artery markings, if present, and apply snugly to the extremity.
- 4. Connect the tubing to the cuff and to the NIBP port on the monitor.
- 5. Rotate the **SPEED DIAL** to outline the NIBP area.
- 6. Press the SPEED DIAL. The NIBP menu appears.
- 7. Select INTERVAL and then select the desired time interval.
- 8. Position the extremity in a relaxed and supported position at approximately the same level as the right atrium of the patient's heart. Inform the patient that the cuff will inflate and cause a "big squeeze" around the arm and that the patient's fingers may tingle.
- Press NIBP to start the measurement, and check that the patient's arm is not moving.
 When the measurement is complete, systolic, diastolic, and mean arterial pressures are displayed. The countdown timer shows the time to the next automatic NIBP measurement.

To cancel a measurement in progress, press NIBP again.

Note: If at any time the cuff pressure exceeds 290 mmHg or there is a system failure of the NIBP module, timer-controlled NIBP is terminated. To reactivate, follow the Timer-Controlled Measurement Procedure.

Cleaning

To clean the cuff and pneumatic tubing:

- 1. Disconnect the tubing from the cuff and monitor. Use a clean, soft cloth dampened with a germicidal solution to wipe clean.
- 2. Inspect the tubing for cracks or kinks. If any damage is noted, replace the tubing.
- 3. Inspect the cuff for damage or excessive wear. If any damage is noted, replace the cuff.
- 4. Allow both to dry before placing the cuff on a patient or reconnecting the tubing to the monitor.

For information about cleaning the device, see Cleaning the Device (on page 212).

Troubleshooting Tips

Table 14 Troubleshooting Tips for NIBP Monitoring

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--------------------------------------|---|---|
| NIBP AIR LEAK message appears | Cuff applied too loosely Leak in cuff/monitor pneumatic system | Check cuff for snug fit on patient. Check that the cuff/monitor connection is secure. Check cuff for leaks. Do not use a cuff that exhibits a leak. |
| NIBP FLOW ERROR message appears | The pneumatic system is not maintaining stable cuff pressure | Deflate or remove cuff.Check tubing for leaks.Replace cuff. |
| NIBP FAILED message appears | The monitor cannot establish zero-pressure reference | Check tubing for kink or blockage. If this message persists, remove monitor from use and obtain service. Use another method to measure the patient's blood pressure. |
| NIBP INITIALIZING message appears | NIBP requested while NIBP module is still initializing | Wait until message disappears and request NIBP. |
| NIBP MOTION message appears | The patient extremity moved too much for the monitor to accurately complete the measurement | Have patient lie quietly with extremity relaxed and supported. Check that patient's arm does not move during NIBP measurement. |
| NIBP OVERPRESSURE message appears | Cuff pressure exceeded 290 mmHg | Disconnect tubing or remove cuff. Avoid very rapid squeezing of the cuff. If this message persists, remove the cuff from use and obtain service. |

Monitoring Noninvasive Blood Pressure

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|--|---|
| NIBP TIME OUT message appears | The monitor did not complete a measurement in 120 seconds | Check cuff for snug fit on patient. Check that cuff artery markings are aligned with the artery. Repeat measurement. Try a higher initial pressure. If this message persists, use another method to measure the patient's blood pressure. |
| NIBP WEAK PULSE message appears | The monitor did not detect any pulses | Check pulses distal to the cuff. Check cuff for snug fit on patient. Check that cuff artery markings are aligned with the artery. |
| XXX appears in place of NIBP readings | NIBP module failed. NIBP module failed to calibrate successfully. | Turn device off and then on again. If problem persists, contact qualified service personnel. |
| NIBP CHECK CUFF message appears | Cuff is not connected to patient or device | Check cuff for snug fit on patient. Check that cuff artery markings are aligned with the artery. Check cuff tubing connection to device. |
| Unable to connect NIBP tubing to device | The LIFEPAK 12 NIBP tubing connector is not compatible with the LIFEPAK 15 NIBP port | Obtain correct NIBP tubing that is compatible with LIFEPAK 15 monitor/defibrill ator. |
| Cuff not deflating | Internal valves fail to open | Disconnect NIBP tubing.Remove cuff from patient. |
| Cuff not inflating | Cuff is not connected to the device | Check tubing connection to device and cuff. |
| | Leak in tubing, cuff, or connector | Replace NIBP tubing or cuff. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Monitoring ETCO2

Intended Use

The end-tidal CO₂ (EtCO₂) monitor is a capnometric device that uses non-dispersive infrared spectroscopy to continuously measure the amount of CO₂ during each breath and report the amount present at the end of exhalation (EtCO₂). The sample is obtained by the side stream method and can be used with intubated or nonintubated patients. Respiration rate is also measured and displayed in breaths per minute.

The EtCO₂ monitor is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times; do not rely solely on the EtCO₂ monitor.

Indications

EtCO₂ monitoring is used to detect trends in the level of expired CO₂. It is used for monitoring breathing efficacy and treatment effectiveness in acute cardiopulmonary care, for example, to determine if adequate compressions are being performed during CPR or to rapidly detect whether an endotracheal tube has been placed successfully.

Contraindications

None known.

EtCO2 Monitoring Warnings

| | 22 Worldoning Warnings | |
|---------|---|--|
| WARNING | Fire Hazard | |
| | Before use, carefully read these operating instructions, the FilterLine® tubing directions for use, and precautionary information. | |
| WARNING | Fire Hazard | |
| | The FilterLine tubing may ignite in the presence of O_2 when directly exposed to laser, electrosurgical devices, or high heat. Use with caution to prevent flammability of the FilterLine tubing. | |
| WARNING | Fire Hazard | |
| | Flammable anesthetics become mixed with the patient's air that is sampled by the capnometer. When using the EtCO ₂ monitor in the presence of flammable gases, such as nitrous oxide or certain other anesthetics, connect the EtCO ₂ gas port to a scavenger system. | |

| WARNING | Possible Inaccurate Patient Assessment |
|----------|---|
| | The EtCO ₂ monitor is intended only as an adjunct in patient assessment and is not to be used as a |
| | diagnostic apnea monitor. An apnea message appears |
| | if a valid breath has not been detected for 30 seconds and indicates the time elapsed since the last valid |
| | breath. It must be used in conjunction with clinical signs |
| | and symptoms. |
| WARNING | Possible Inaccurate CO₂ Readings |
| WARNING | • |
| | Using other manufacturers' CO₂ accessories may cause the device to perform improperly and invalidate the |
| | safety agency certifications. Use only the accessories |
| | that are specified in these operating instructions. |
| | Possible Strangulation |
| WARNING | 1 033ibic Ottangulation |
| | Carefully route the patient tubing (FilterLine) to reduce the possibility of patient entanglement or strangulation. |
| WARNING | Infection Hazard |
| WARINING | |
| | Do not reuse, sterilize, or clean Microstream® CO₂ accessories as they are designed for single-patient |
| | |
| | one-time use. |
| WARNING | , |
| WARNING | one-time use. |

How Capnography Works

An EtCO₂ sensor continuously monitors carbon dioxide (CO₂) that is inspired and exhaled by the patient. The sensor employs Microstream non-dispersive infrared (IR) spectroscopy to measure the concentration of CO₂ molecules that absorb infrared light.

The CO_2 FilterLine system delivers a sample of the exhaled gases directly from the patient into the LIFEPAK 15 monitor for CO_2 measurement. The low sampling flow rate (50 ml/min) reduces liquid and secretion accumulation and prevents obstruction, which maintains the shape of the CO_2 waveform.

The CO₂ sensor captures a micro sample (15 microliters). This extremely small volume allows for fast rise time and accurate CO₂ readings, even at high respiration rates.

The Microbeam IR source illuminates the sample cell and the reference cell. This proprietary IR light source generates only the specific wavelengths characteristic of the CO₂ absorption spectrum. Therefore, no compensations are required when concentrations of O₂, anesthetic agent, or water vapor are present in the exhaled breath.

You can set up the LIFEPAK 15 monitor/defibrillator to use the capnography Body Temperature Pressure Saturated (BTPS) conversion method. This option corrects for the difference in temperature and moisture between the sampling site and alveoli. The correction formula is 0.97 × the measured EtCO₂ value. See the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

EtCO2 Monitoring Waveform Analysis

Valuable information concerning the patient's expired CO₂ can be acquired by examination and interpretation of the waveform.

The Phases of the Waveform

The following figure is a graphic representation of a normal capnograph waveform. Four phases of the waveform require analysis. The flat I–II baseline segment (Respiratory Baseline) represents continued inhalation of CO₂-free gas. This value normally is zero. The II–III segment (Expiratory Upstroke), a sharp rise, represents exhalation of a mixture of dead space gases and alveolar gases from acini with the shortest transit times. Phase III–IV (Expiratory Plateau) represents the alveolar plateau, characterized by exhalation of mostly alveolar gas. Point IV is the end-tidal (EtCO₂) value that is recorded and displayed by the monitor. Phase IV–V (Inspiratory Downstroke), a sharp fall, reflects the inhalation of gases that are CO₂-free. Alterations of the normal capnograph or EtCO₂ values are the result of changes in metabolism, circulation, ventilation, or equipment function.

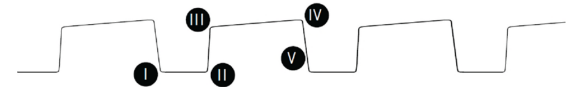


Figure 27 Phases of the Respiratory Waveform

Respiratory Baseline Elevation of the waveform baseline (I–II segment) usually represents rebreathing CO₂. This elevation usually is accompanied by gradual increases in the EtCO₂ value. Rebreathing CO₂ is common in circumstances of artificially produced increased dead space and hypoventilation. Precipitous rises in both baseline and EtCO₂ values usually indicate contamination of the sensor.

Expiratory Upstroke In the normal waveform, the rising phase (II–III segment) is usually steep. When this segment becomes less steep, CO₂ delivery is delayed from the lungs to the sampling site. The causes of this delay can be physiologic or mechanical and include bronchospasm, obstruction of the upper airway, or obstruction (or kinking) of an endotracheal tube (ETT).

Expiratory Plateau The plateau of the waveform, which represents the remainder of expiration (III-IV segment), should be nearly horizontal. The end of the plateau represents the EtCO₂ value. Upward slanting of the expiratory plateau occurs when there is uneven emptying of the alveoli. Similar to the diminished slope of the Expiratory Upstroke, this pattern can occur in asthma, chronic obstructive pulmonary disease (COPD), partial upper-airway obstruction, or partial mechanical obstruction such as a partially kinked ETT.

Monitoring ETCO2

Inspiratory Downstroke The fall to baseline (IV-V segment) is a nearly vertical drop. This slope can be prolonged and can blend with the expiratory plateau in cases of leakage in the exhale portion of the breathing circuit. The peak EtCO₂ value (IV) is often not reached. Relying on the numeric end-tidal value without observing the breathing waveform may obscure the presence of a leak.

EtCO2 Monitoring Procedure

When activated, the EtCO₂ monitor draws power from the defibrillator. The LIFEPAK 15 monitor/defibrillator activates the EtCO₂ monitor when it senses the attachment of the FilterLine set. Initialization, self-test, and warm up of the EtCO₂ monitor is typically less than 30 seconds, but may take up to two-and-one-half minutes.

CAUTION

Possible Equipment Damage

Failure to replace a broken or missing CO_2 port door may allow water or particulate contamination of the internal CO_2 sensor. This may cause the CO_2 module to malfunction.

To monitor EtCO₂:

- 1. Press ON.
- 2. Select the appropriate EtCO₂ accessory for the patient.
- 3. Open the CO₂ port door and insert the FilterLine connector; turn connector clockwise until tight.
- 4. Verify that the CO₂ area is displayed. The EtCO₂ monitor performs the autozero routine as part of the initialization self-test.

Note: If you use a ventilation system, do not connect the FilterLine set to the patient/ventilation system until the EtCO₂ monitor has completed its self-test and warm-up.

- Display CO₂ waveform in Channel 2 or 3.
- 6. Connect the CO₂ FilterLine set to the patient.
- Confirm that the EtCO₂ value and waveform are displayed. The monitor automatically selects
 the scale for the best visualization of the waveform. You can change the scale, if desired, as
 described in the next section.

Note: It is possible for the FilterLine set to become loose at the device connection and still have an EtCO₂ value and CO₂ waveform, but they may be erroneously low. Make sure the FilterLine connection is firmly seated and tight.

Note: The capnography module performs self-maintenance within the first hour of monitoring and once an hour during continuous monitoring. The self-maintenance includes "auto-zeroing." Self-maintenance is also initiated when the surrounding temperature changes 8°C (14.4°F) or more, or the surrounding pressure changes greater than 20 mmHg. The CO₂ module detects this change and attempts to purge the tubing. To clear the **CO2 FILTERLINE PURGING** or **CO2 FILTERLINE BLOCKAGE** messages, remove the FilterLine tubing and reconnect it to the monitor.

CO2 Display

The following scales are available to display the CO₂ waveform. The LIFEPAK 15 monitor/defibrillator automatically selects the scale based on the measured EtCO₂ value. To change the CO₂ scale, outline and select the CO₂ area using the **SPEED DIAL** and then select the desired scale from the scale menu.

- Autoscale (default)
- 0-20 mmHg (0-4 Vol% or kPa)
- 0-50 mmHg (0-7 Vol% or kPa)
- 0-100 mmHg (0-14 Vol% or kPa)

The CO₂ waveform is compressed (displayed at 12.5 mm/sec sweep speed) to provide more data in the 4-second screen. There is a slight delay between when the breath occurs and when it appears on the screen. Printouts are at 25 mm/sec. Continuous print may be changed to 12.5 mm/sec, if desired.

The monitor shows the maximum CO₂ value over the last 20 seconds. If the EtCO₂ values are increasing, the change can be seen with every breath. However, if the values are continually decreasing, it will take up to 20 seconds for a lower numerical value to be displayed. Because of this, the EtCO₂ value may not always match the level of the CO₂ waveform.

CO₂ Alarms

The EtCO₂ monitor provides:

- EtCO₂ high and low alarms controlled by activating ALARMS (see Alarms (on page 39))
- FiCO₂ (inspired CO₂) alarm (automatic and not adjustable)
- Apnea alarm (automatic and not adjustable)

Note: The apnea alarm occurs when a breath has not been detected for 30 seconds. The message **ALARM APNEA** appears in the message area along with the time since the last detected breath.

CO2 Detection

A CO_2 waveform appears when any CO_2 is detected, but CO_2 must be greater than 3.5 mmHg for a numerical value to be displayed. However, the CO_2 module will not recognize a breath until the CO_2 is at least 8 mmHg (1.0% or kPa). Valid breaths must be detected in order for the apnea alarm to function and to count the respiratory rate (RR). The RR represents an average over the last eight breaths.

Monitoring ETCO2

When CO₂ is not detected in the cardiac arrest situation—for example, the CO₂ waveform is either dashes "---" or a flat solid line at or near zero—several factors must be quickly evaluated. Assess for the following causes:

Equipment issues

- Disconnection of the FilterLine set from the endotracheal tube (ETT)
- System is purging due to fluid in the patient/sensor connection from ET administration of medications
- · System is auto-zeroing
- Shock was delivered and system is resetting
- Loose FilterLine set to device connection

Loss of airway function

- Improper placement of ETT
- ETT dislodgment
- ETT obstruction

Physiological factors

Apnea

- Loss of perfusion
- Exsanguination

- Massive pulmonary embolism
- Inadequate CPR

Cleaning

Accessories for CO₂ monitoring are disposable and are intended for single-patient use. Do not clean and reuse a FilterLine set. Dispose of the contaminated waste according to local protocols.

For information about cleaning the device, see Cleaning the Device (on page 212).

Troubleshooting Tips

Table 15 Troubleshooting Tips for EtCO2 Monitoring

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|---|---|
| ALARM APNEA message appears and waveform is solid line at or near zero | No breath has been detected for 30 seconds since last valid breath | Check the patient. |
| | FilterLine connection to device is loose | Twist FilterLine connector clockwise until tight and firmly seated. |
| | FilterLine set is disconnected from patient or ETT | Check ventilation equipment (if used) for leaks or disconnected tubing. |
| CO2 FILTERLINE OFF message appears and waveform is "" | FilterLine set disconnected or not securely connected to device | Connect FilterLine set to device port. Twist FilterLine connector clockwise until tight and firmly seated. |
| CO2 FILTERLINE PURGING message appears and waveform is "" | FilterLine set is kinked or clogged with fluid, or rapid altitude change occurred | Disconnect and then reconnect the FilterLine set. Twist FilterLine connector clockwise until tight and firmly seated. |
| CO2 FILTERLINE BLOCKAGE message appears and waveform is "" | The message appears after 30 seconds of unsuccessful purging | Disconnect and then reconnect the FilterLine set.Change the FilterLine set. |
| | FilterLine set is kinked or clogged | Twist FilterLine connector clockwise until tight and firmly seated |
| CO2 INITIALIZING message appears and waveform is "" | FilterLine set connected to device while module is initializing | None. |
| | Defibrillation shock delivered | None. System resets automatically within 20 seconds. |
| AUTO ZEROING message appears and waveform is "" | Module is performing self- maintenance | None. |
| | Defibrillation shock delivered | None. System resets automatically within 20 seconds. |
| EtCO ₂ values are erratic | FilterLine connection to device is loose | Twist FilterLine connector clockwise until tight and firmly seated. |
| | A leak in the FilterLine set | Check for connection leaks and line leaks to patient, and correct, if necessary. |
| | A mechanically ventilated patient breathes spontaneously or patient is talking | No action required. |

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Monitoring ETCO2

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|--|---|
| EtCO ₂ values are consistently higher than expected | Physiological cause such as COPD | None. |
| | Inadequate ventilation | Check ventilator, increase ventilatory rate/bagging. |
| | Patient splinting during breathing | Supporting measures such as pain relief. |
| | Improper calibration | Contact qualified service personnel. |
| EtCO ₂ values are consistently lower than expected | FilterLine connection to device is loose | Twist FilterLine connector clockwise until tight and firmly seated |
| | Physiological cause | See Physiological factors in CO2 Detection (on page 91). |
| | Hyperventilation | Check ventilator, decrease ventilatory rate/bagging. |
| | Improper calibration | Contact qualified service personnel. |
| CO ₂ waveform stays elevated for several seconds | Expiration is prolonged due to bagging technique | Release bag reservoir completely with expiration. Observe that elevated baseline returns to normal level. |
| Sudden extreme increase in EtCO ₂ | Fluid has entered CO ₂ module | Contact qualified service personnel. |
| XXX appears instead of EtCO ₂ value | CO₂ module malfunction | Turn device off and then on again. |
| | | If problem persists, contact qualified service personnel. |
| There is no EtCO ₂ value and the CO ₂ waveform is flat | Measured CO ₂ is less than 3.5 mmHg | See CO2 Detection (on page 91). |

Note: To decrease the likelihood of the FilterLine connection coming loose during use, handstraighten the tubing after removal from the package before connecting to patient or device.

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Monitoring Invasive Pressure

Intended Use

The LIFEPAK 15 invasive pressure (IP) monitor is intended for measuring arterial, venous, intracranial, and other physiological pressures using an invasive catheter system with a compatible transducer.

The IP monitor is a tool to be used in addition to patient assessment. Care should be taken to assess the patient at all times; do not rely solely on the IP monitor.

Indications

Invasive pressure monitoring is indicated for use in patients who require continuous monitoring of physiological pressures in order to rapidly assess changes in the patient's condition or response to therapy. It may also be used to aid in medical diagnosis.

Contraindications

None known.

IP Monitoring Warnings

| W | | | |
|---|--|--|--|
| | | | |
| | | | |

Possible Inaccurate Pressure Readings, Air Embolism, Blood Loss, or Loss of Sterility

Before use, carefully read these operating instructions, and the transducer and infusion set instructions for use and precautionary information.

WARNING

Inaccurate Pressure Readings

Pressure readings should correlate with the patient's clinical presentation. If readings do not correlate, verify that the zeroing stopcock is positioned at the patient's zero reference, rezero the transducer, and/or check the transducer with a known or calibrated pressure. Manually check cuff blood pressure.

WARNING

Inaccurate Pressure Readings

Changing the patient's position changes the zero reference level. Relevel the transducer's zeroing stopcock any time the patient's position is changed.

Monitoring Invasive Pressure

WARNING

Possible Patient Injury or Equipment Damage

Use only IP transducers that are specified for use with this device. Protection of the device against defibrillator discharge is dependent on the use of IP transducers that are specified by Physio-Control.

WARNING

Possible Lethal Arrhythmia

Ventricular fibrillation may be induced if the isoelectric barrier of the transducer is disrupted. The isoelectric barrier within the transducer may be disrupted if the transducer body is damaged. Do not use a transducer that is visibly damaged or leaking fluid.

WARNING

Increased Intracranial Pressure

Do not use a continuous flush device with transducers used for intracranial monitoring.

IP Monitoring

Two channels are available for invasive pressure monitoring, with default labels P1 and P2 and the user-selectable labels shown in the following table.

Table 16 IP Labels and Descriptions

| LABEL | DESCRIPTION | |
|-------|---------------------------|--|
| ART | Arterial Pressure | |
| PA | Pulmonary Artery Pressure | |
| CVP | Central Venous Pressure | |
| ICP | Intracranial Pressure | |
| LAP | Left Atrial Pressure | |

When the default labels P1 and P2 are used, the IP monitoring area displays systolic, diastolic, and mean pressures. When ICP, LAP, or CVP labels are used, the IP monitoring area displays mean pressure in large type. Systolic and diastolic pressures are not displayed.

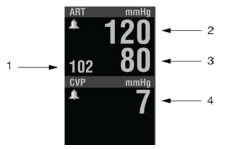


Figure 28 IP Labels

FIGURE LEGEND

- 1 ART mean pressure
- 2 ART systolic pressure
- 3 ART diastolic pressure
- 4 CVP mean pressure

Because pressures can change in a short time, data should be checked regularly during vital sign monitoring.

How IP Monitoring Works

IP monitoring involves the conversion of fluid pressure into an electrical signal. The conversion is accomplished with a pressure transducer. The transducer is connected to a patient's indwelling pressure catheter using a special assembly of tubing, stopcocks, adapters, flush valves, and fluids, commonly known as a flush system. The transducer translates the pressure wave into an electrical signal. A well-functioning flush system is essential for obtaining undistorted waveforms and accurate information.

IP monitoring is available on either Channel 2 or 3. The IP monitor is compatible with industry standard (IEC 60601-2-34) pressure transducers with $5\mu V/V/mmHg$ sensitivity. The transducer must provide defibrillation protection of at least 360 joules. The following IP transducers may be used with the LIFEPAK 15 monitor/defibrillator.

| MANUFACTURER | DESCRIPTION |
|----------------------|---|
| Utah Medical | DPT-100 Deltran® Disposable Pressure Transducer |
| Edwards Lifesciences | TruWave™ Disposable Pressure Transducer |
| ICU Medical | Transpac® IV Disposable Pressure Transducer |

An invasive pressure adapter cable is used to connect the transducer to the monitor. The IP connector is a 6-pin type 3102A-14S-6S connector. The connector pinout has the following configuration, counterclockwise from 12 o'clock, viewed from the front of the LIFEPAK 15 monitor/defibrillator.

| A pin = - signal | B pin = + excitation | C pin = + signal |
|----------------------|----------------------|-------------------|
| D pin = - excitation | E pin = shield | F pin = unlabeled |

IP Monitoring Procedure

Prepare a flush system according to local protocols. Position the transducer at the patient's phlebostatic axis (zero-reference level).

To avoid offset errors, a zero reference must be established before any meaningful pressure readings are obtained. This is done by opening the transducer stopcock to air so that atmospheric pressure becomes the reference.

The P1 or P2 connector and Channel 2 or 3 can be used for IP monitoring. P1 and Channel 2 are used in these instructions.

To monitor IP:

- 1. Prepare the transducer system according to the operating instructions provided with the transducer and your local protocol.
- 2. Press ON.
- 3. Connect the IP cable to the transducer and to the P1 port on the monitor.
- 4. Use the default label **P1** or select **ART**, **PA**, **CVP**, **ICP**, or **LAP**. To change the label, select the P1 area. From the menu, select **P1**. Select a label from the list.
- Use the SPEED DIAL to outline and select CHANNEL 2 on the Home Screen. From the Channel 2 menu, select WAVEFORM and then select the label that is desired for the waveform.
- 6. Open the transducer's stopcock to air to zero the transducer and remove stopcock cap. Select the **P1** area. Select **ZERO** from the menu. The message **P1 ZEROED** appears when zeroing is complete and the pressure values are displayed as zeros.
- Close the stopcock to air. The patient's pressure waveform should be displayed. A scale is automatically selected to display the pressure. Confirm that pressure amplitude correlates with the digital readout.

Note: If you place a cap on an open port before you close the port to air, an error message may appear. You will be required to zero the transducer again.

If pressure alarms are desired, set the alarms after you obtain a satisfactory waveform. Error or alarm messages appear in the message area at the bottom of the screen. For more information, see Alarms (on page 39).

IP Scale Options

The IP monitor can display pressures from -30 to 300 mmHg. After zeroing the transducer pressure, the monitor automatically selects one of the following scales based on the patient's measured pressure:

- -30 to 30 mmHg
- 0 to 60 mmHg
- 0 to 120 mmHg
- 0 to 150 mmHg
- 0 to 180 mmHg
- 0 to 300 mmHg

You can also manually select one of these scales or autoscale to readjust the waveform within the channel.

To change the scale:

- 1. Use the SPEED DIAL to outline and select the P1 area. The P1 menu appears.
- 2. From the menu, select SCALE and then choose a scale from the list.

Cleaning and Inspection

IP transducers are disposable and are intended for single-patient use. Do not clean and reuse transducers. Dispose of the contaminated waste according to local protocols.

IP cables are reusable and may be cleaned. To clean the reusable IP cable:

- 1. Disconnect the cable from the monitor.
- 2. Inspect the cable for damage or wear.
- 3. Use a clean, soft cloth dampened with a germicidal solution to wipe clean.
- 4. Allow to dry before reconnecting the cable to the monitor.

For information about cleaning the device, see Cleaning the Device (on page 212).

Troubleshooting Tips

The error messages in the following table use the text **PX** to represent any of the labels for invasive pressure, including P1, P2, and the user-selectable labels ART, PA, CVP, ICP, and LAP.

Table 17 Troubleshooting Tips for IP Monitoring

| Table 17 Troubleshooting Tips for | | CORRECTIVE ACTION |
|---------------------------------------|---|--|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
| Invasive pressure value is blank | No transducer is connected | Connect the transducer to the cable, and the cable to the monitor. |
| No scale appears next to the waveform | The zero reference has not been established | Zero the transducer. |
| PX NOT ZEROED message appears | The zero reference has not been established | Zero the transducer. |
| PX ZERO FAILED message appears | An unsuccessful attempt has been made to set a zero reference value | Make sure that the transducer is open to air and repeat the attempt to zero. |
| Dampened waveform | Loose connection | Check the entire system for leaks. Tighten all connections. Replace any defective stopcocks. |
| | Tubing too long or too compliant | Use short, stiff tubing with large diameter. |
| | Thrombus formation, air bubbles, or blood left in catheter after blood draw | Use syringe to draw back air or particles in catheter, and then flush system. |
| | Kinked catheter, catheter tip against vessel wall, arterial spasm | Reposition catheter. Anchor catheter to skin at insertion site. |
| Resonating waveform | Tubing too long | Use short, stiff tubing with a large diameter. |
| No waveform. No pressure reading. | Transducer closed to patient | Check patient. Check stopcock positions and monitor setup. |
| | Defibrillator shock just delivered | None. |
| Invasive BP lower than cuff BP | Transducer level higher than the heart | Reposition transducer to correct height. |
| | Loose connection | Tighten all connections. |
| | Thrombus formation, air bubbles, or blood in catheter, kinking, or arteriospasm | Use syringe to draw back air or particles in catheter, and then flush system. |
| | Improper zero reference | Open stopcock to air and rezero transducer. |
| | Defective transducer | Replace transducer. |

Chapter 4 | Monitoring

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|--|--|
| Invasive BP higher than cuff BP | Transducer level lower than the heart | Reposition transducer to correct height. |
| | Improper zero reference | Rezero. |
| | Catheter whip artifact | Change catheter tip position. Use mean pressure values (mean pressure is less affected by extremes and will therefore reflect a more accurate reading). |
| Inability to flush system | Pressure bag leaking | Keep positive pressure in flush bag at all times. Remove dressing to check for external kinking. |
| | Partially kinked or obstructed catheter | Replace catheter, if clotted. |
| Inability to zero system | Stopcock not open to air or defective | Check stopcock position. Replace any defective stopcocks. |
| | Defective transducer | Replace transducer. |
| System has been zeroed but continues to indicate zero reference required | Steps to zero system performed in wrong order | Close stopcock to air before placing cap on port. |
| Catheter whip (fling) artifact Pulmonary Artery | Excessive catheter movement. Motion of the catheter tip within the vessel accelerates fluid movement in the catheter, causing artifact to be superimposed on the pressure wave, increasing readings by 10–20 mmHg. | Change catheter tip position. Use mean pressure values (mean pressure is less affected by extremes and therefore reflects a more accurate reading). |
| Permanent Pulmonary Wedge Pressure (PWP) tracing (wedge tracing persists after balloon deflation) | Catheter tip partially clotted | Use syringe to aspirate, and then flush. |
| | Catheter migrated distally in pulmonary artery | Observe PA waveform before balloon inflation. Flattening of the waveform could indicate wedging with balloon deflated. Turn patient side to side in Trendelenburg position, or stimulate cough in attempt to dislodge catheter. Retract catheter with balloon deflated until proper position is obtained. Minimize chances of catheter advancement by firmly anchoring catheter at insertion site. |

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Monitoring Invasive Pressure

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|------------------------------|--|--|
| Failure to obtain PWP | Malposition of catheter tip | Reposition catheter. |
| | Leak in balloon. Ruptured balloon. | Replace catheter. |
| Progressive elevation of PWP | Overinflation | Inflate balloon in small increments while watching scope for confirmation of wedging. Use only enough air to wedge. Do not use more than the volume recommended by the manufacturer. |
| | Catheter migrated distally in pulmonary artery | Reposition catheter. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Monitoring Continuous Temperature

Intended Use

The LIFEPAK 15 temperature monitor is intended for continuous monitoring of body temperature.

Indications

Temperature monitoring is indicated for use in patients who require continuous monitoring of body temperature.

Contraindications

None known.

Temperature Monitoring Warnings

| W | | | |
|---|--|--|--|
| | | | |

Possible Inaccurate Temperature Readings

Using temperature probes or cables that are not approved by Physio-Control may cause improper temperature monitoring performance and invalidate safety agency certifications. Use only probes and cables that are specified in these operating instructions.

WARNING

Possible Inaccurate Temperature Readings

The Measurement Specialties 4400 Series temperature probes must be used with the adapter cable that is listed on the Physio-Control website. Using other manufacturers' connector cables may cause the device to perform improperly.

WARNING

Infection Hazard

The temperature probe is disposable and intended for single-patient use. Do not clean and reuse temperature probes. Dispose of contaminated waste according to local protocols.

WARNING

Possible Strangulation

Carefully route the temperature probe cable to reduce the possibility of patient entanglement or strangulation.

How Temperature Monitoring Works

The temperature probe contains a thermistor which converts temperature to electrical resistance. The LIFEPAK 15 monitor/defibrillator measures the resistance and converts it into degrees Celsius or Fahrenheit. The probe accuracy is ±0.1°C.

Note: Celsius or Fahrenheit reporting may be selected in Setup mode. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

The temperature area of the home screen is blank until a temperature value between 24.8° and 45.2°C (76.6° and 113.4°F) is detected. When a temperature value in this range is detected, the value is automatically displayed.

After a valid body temperature between 31° and 41°C (87.8° and105.8°F) is detected, the device monitors the temperature value for possible sensor dislodgement or disconnection. If the device detects a temperature outside of the valid body temperature range, the **TEMP: CHECK SENSOR** message appears. The following table shows the screen messages and temperature values that are displayed for each temperature range.

Table 18 Temperature Values and Messages

| TEMPERATURE | MESSAGE | TEMP VALUE DISPLAY |
|-----------------------------------|--------------------------|--------------------|
| Less than 24.8°C (76.6°F) | TEMP: CHECK SENSOR | Dashes () |
| 24.8° to 30.9°C (76.6° to 87.6°F) | TEMP: CHECK SENSOR | Current temp value |
| 31° to 41°C (87.8° to 105.8°F) | No message (valid range) | Current temp value |
| 41.1° to 45.2°C (106° to 113.4°F) | TEMP: CHECK SENSOR | Current temp value |
| Greater than 45.2°C (113.4°F) | TEMP: CHECK SENSOR | Dashes () |
| Temperature probe disconnected | TEMP: CHECK SENSOR | Dashes () |

The temperature monitor performs an accuracy check each time it is turned on, and periodically while monitoring temperature. If the temperature accuracy check fails, the message **TEMP: ACCURACY OUTSIDE LIMITS** is displayed, and the temperature value is "XXX".

Temperature Monitoring Equipment

The following accessories are required for temperature monitoring:

- Temperature adapter cable
- Measurement Specialties 4400 Series disposable temperature probe. You can use the following probe types with the LIFEPAK 15 monitor/defibrillator:
 - Esophageal/rectal
 - Foley catheter
 - Skin (Note: Measurement Specialties skin temperature probe 4499HD is approved for use with the LIFEPAK 15 monitor/defibrillator. Do not use Measurement Specialties part number 4499.)

For a list of the accessories that are intended for use with the LIFEPAK 15 monitor/defibrillator, contact your Physio-Control representative or see the LIFEPAK 15 Monitor/Defibrillator Accessory Catalog at www.physio-control.com. Carefully read the Instructions for Use that are

provided with the probes and connector cable for sensor placement instructions, use instructions, warnings, cautions, and specifications.

IMPORTANT! The Instructions for Use that are provided with the Measurement Specialties temperature probes refer to a connector cable that is not compatible with the LIFEPAK 15 monitor/defibrillator. Only use the adapter cable that is approved for use with the LIFEPAK 15 monitor/defibrillator.

Temperature Monitoring Procedure

- 1. Connect the temperature adapter cable to the TEMP port on the monitor/defibrillator.
- 2. Connect the temperature probe to the temperature adapter cable.
- 3. Attach the temperature probe to the patient as described in the temperature probe Instructions for Use.

Notes:

- The temperature area on the display is not activated until the monitor/defibrillator detects a temperature between 24.8° and 45.2°C (76.6° and 113.4°F). To manually activate the temperature monitoring area, use the **SPEED DIAL** to outline and select the temperature area on the Home Screen. From the menu, select **ON**.
- The temperature probe may require 3 minutes to equilibrate after placement on the patient monitoring site.
- 4. Confirm that the temperature reading appears and is stable.
- Use the default label TEMP or select one of the user-selectable labels shown in the following table. To change the label, select the TEMP area. From the menu, select TEMP. Select a label from the list.

Table 19 TEMP Labels and Descriptions

| LABEL | DESCRIPTION |
|-----------|----------------------------|
| T-esoph | Esophageal Temperature |
| T-naso | Nasopharangeal Temperature |
| T-bladder | Bladder Temperature |
| T-rectal | Rectal Temperature |
| T-skin | Skin Temperature |

Cleaning and Disposal

Temperature probes are disposable and intended for single-patient use. Do not clean and reuse temperature probes. Dispose of the contaminated waste according to local protocols.

Temperature adapter cables are reusable and may be cleaned. To clean the reusable temperature cable:

- 1. Disconnect the cable from the monitor.
- 2. Use a clean, soft cloth dampened with a germicidal solution to wipe clean. See Cleaning the Device (on page 212) for a list of acceptable cleaning solutions.
- 3. Allow to dry before reconnecting the cable to the monitor.

For information about cleaning the device, see Cleaning the Device (on page 212).

Troubleshooting Tips

Table 20 Troubleshooting Tips for Temperature Monitoring

| Table 20 Troubleshooting Tips for Temperature Monitoring | | | | |
|--|---|--|--|--|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION | | |
| CHECK SENSOR message appears and value is "" | Temperature value is out of range | Check that probe is positioned properly. | | |
| | Temperature probe is dislodged or positioned incorrectly | Check that probe is positioned properly. | | |
| | Probe not connected to cable, or cable not connected to device | Check that probe and cable are connected properly. | | |
| | Damaged cable or probe | Replace damaged cable or probe. | | |
| CHECK SENSOR message appears while value is displayed | Temperature probe is dislodged and value is below 31°C (87.8°F) | Check that probe is positioned properly. | | |
| | Temperature probe is dislodged and value is above 41.0°C (105.8°F) | Check that probe is positioned properly. | | |
| TEMP: ACCURACY OUTSIDE LIMITS message appears and value is XXX | Temperature accuracy check failed | Turn device off and then on again.If problem persists, contact qualified service personnel. | | |
| XXX appears in place of temperature reading | Temperature module is not calibrated | Turn device off and then on again.If problem persists, contact qualified service personnel. | | |
| | Temperature module failed | Turn device off and then on again.If problem persists, contact qualified service personnel. | | |

Chapter 4 | Monitoring

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|---|---|
| Temperature area of home screen is blank | Initial temperature not automatically displayed until device detects temperature between 24.8° and 45.2°C (76.6° and 113.4°F) | Allow up to 3 minutes for probe to equilibrate. Check that probe is positioned properly. |
| | Temperature probe not detected by device | Check connections between probe, adapter cable, and device. |
| | | Check that the sensor is approved for use with the LIFEPAK 15 monitor/defibrillator. |
| | | Contact qualified service personnel. |

Vital Sign and ST Segment Trends

Intended Use

The trends feature of the LIFEPAK 15 monitor/defibrillator provides the ability to graphically display and document the patient's vital signs (VS) and ST segment measurements for up to eight hours. VS trending is intended for use with any patient who requires continuous monitoring of vital signs over an extended period of time to identify changes in patient condition and to document patient response to therapy. ST trending is intended for use with patients suspected of having acute ischemic events, such as unstable angina, and for patients during treatment of an acute ischemic event. ST segment measurement is initiated using a 12-lead ECG and is derived using the University of Glasgow 12-Lead ECG Analysis Program.

VS and ST Trends Warning

WARNING

Inaccurate Interpretation of Patient Status

Vital sign and ST graphs are tools to be used in addition to patient assessment. Artifact and noise may produce spurious data. Ensure artifact-free monitoring as much as possible and assess the patient frequently to confirm the appropriateness of monitor data.

How VS Trends Work

Each active vital sign can be displayed graphically for time ranges of 30 minutes, and 1, 2, 4, and 8 hours. The vital signs are HR, SpO₂, SpCO, SpMet, CO₂, Temp, and RR; and systolic, diastolic, and mean pressures. Data is sampled every 30 seconds. If valid data is not available, a blank space is substituted on the graph. NIBP values are plotted only when an NIBP measurement is obtained. VS measurements are not averaged or filtered. No messages or alarms occur based on changes in VS measurements.

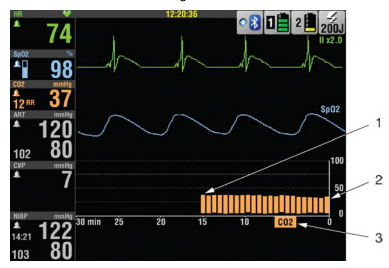


Figure 29 EtCO2 Trend Graph

FIGURE LEGEND

- 1 First EtCO₂ measurement
- 2 Most recent EtCO₂ measurement
- 3 VS label

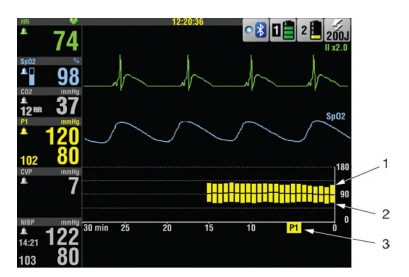


Figure 30 Pressure Trend Graph

FIGURE LEGEND

- 1 Systole pressure
- 2 Diastolic pressure
- 3 VS label

How ST Trends Work

ST measurements can be displayed graphically for time ranges of 30 minutes, and 1, 2, 4, and 8 hours. ST trending is initiated by obtaining the patient's first 12-lead ECG. The ST J-point (STJ) is the part of the ST segment that is measured (see the following figure). The STJ measurement is plotted on the ST trend graph (see Figure, ST Trend Graph (on page 110)).



Figure 31 STJ Measurement

When all leads of the 12-lead ECG cable are attached to the patient, STJ measurements are obtained automatically every 30 seconds. If a lead is off, or the ECG data is too noisy, ST measurements are not obtained and the graph shows a blank for that time period. If an STJ measurement in any lead deviates from the initial measurement by 1 mm (0.1 mV) or more and the deviation persists for 2.5 minutes, the monitor automatically prints another 12-lead ECG. Manual requests for 12-lead ECGs do not affect ST trending or automatic printing.

Interpreting the ST Trend Graph

Using the first 12-lead ECG, the monitor identifies the presence of any STJ displacement, either negative or positive, and the lead that has the most STJ displacement. When **AUTO** is selected, the lead that has the most STJ displacement is shown on the graph. The STJ is measured every 30 seconds thereafter.

The following figure shows an example of an ST trend graph. The elapsed time goes from right to left across the screen. The most current STJ measurement is on the far right. Each time an STJ measurement is obtained, it is compared to the first STJ or baseline measurement. The bars represent the change in the STJ compared to the first measurement.

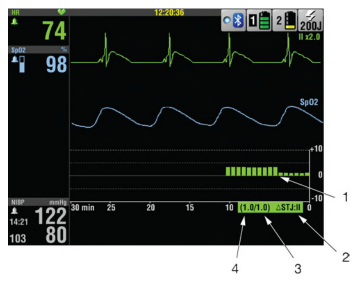


Figure 32 ST Trend Graph

FIGURE LEGEND

- 1 Increase and then decrease in STJ
- 2 Lead
- 3 Change in STJ
- 4 Current STJ

This ST trend graph depicts the changes in STJ from a patient's first 12-lead ECG over 10 minutes of monitoring time. The patient's initial ECG showed no ST elevation in any lead. Then the patient developed 3 mm elevation in Lead II. This change in ST elevation is represented by the vertical bars and lasted approximately 5 minutes. (Each vertical bar represents a 30-second interval). After treatment was initiated, the ST decreased to the current STJ measurement of 1.0, but is still positive compared to the initial ECG.

The annotation (1.0/1.0) means that the current STJ measurement is elevated 1.0 mm and represents a change of 1.0 mm from the initial ECG. To confirm the value of the initial 12-lead ECG STJ measurement, subtract the STJ change from the current STJ measurement, for example, 1.0 - 1.0 = 0. You can display the ST graph of other leads.

Displaying and Printing Trend Graphs

The trend graph for any active vital sign or ST measurement can be displayed in Channel 2 or 3. The example in Figure, ST Trend Graph (on page 110), shows the trend graph in Channel 3. Only two trend graphs can be displayed at a time, but the device collects trend data on all active vital sign values.

To display trend graphs:

- 1. Rotate the **SPEED DIAL** to outline Channel 2 or 3, and then press the **SPEED DIAL** to select the channel. The Channel menu appears.
- Select WAVEFORM, and then select TREND.
- 3. Select **SOURCE**, and then select the desired VS or ST.
- 4. The default setting for SCALE and RANGE is AUTO. When AUTO is used, the monitor automatically updates the scale so that all values are displayed and all data from Power On to the present time is visible. If you change scale or range, some data may not be visible because it is off scale or out of range.
- 5. Press **HOME SCREEN**. The graph for the selected VS or ST appears in the channel.

Note: To initiate ST trends, you must obtain a 12-lead ECG. The initial ECG provides the baseline ST measurement and initiates the ST trends feature.

To print trend graphs:

- 1. Press **OPTIONS**. The Options menu appears.
- Rotate and then press the SPEED DIAL to select PRINT.
- 3. Select **REPORT**, and then select **TREND SUMMARY**.
- Select PRINT. The Trend Summary Report prints graphs of all actively monitored VS and ST trends.

VS and ST Monitoring Considerations

For best results, consider the following:

- The ability of the patient to cooperate and be relaxed. Patients who are restless can produce noisy physiological signals. Noisy signals can result in inaccurately high or low data measurements.
- The quality of the physiological signal. If the ECG has significant artifact, the HR may have spurious measurements. Noisy 12-lead ECGs may need to be overridden, and ST measurements will not be obtained.
- The expected length of time the patient is to be monitored. VS graphs of the patient monitored for only a short time (for example, 15 minutes) may not provide enough data to identify gradual changes in patient condition.
- The patient ECG rhythm. Diagnosis of ST associated ischemia is inhibited by certain ECG findings such as left bundle branch block and ventricular pacing.

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Chapter 5

Therapy

This chapter describes patient therapy.

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|---|-----|
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| Manual Defibrillation | 132 |
| Synchronized Cardioversion Procedure | 137 |
| Noninvasive Pacing | 141 |
| Pediatric ECG Monitoring and Manual Mode Therapy Procedures | 147 |

General Therapy Warnings and Cautions

WARNING

Shock Hazard

The defibrillator delivers up to 360 joules of electrical energy. When discharging the defibrillator, do not touch the paddle electrode surfaces or disposable therapy electrodes.

WARNING

Shock Hazard

If a person is touching the patient, bed, or any conductive material in contact with the patient during defibrillation, the delivered energy may be partially discharged through that person. Clear everyone away from contact with the patient, bed, and other conductive material before discharging the defibrillator.

WARNING

Shock Hazard

Do not discharge the defibrillator into the open air. To remove an unwanted charge, change the energy selection, select disarm, or turn off the defibrillator.

WARNING

Possible Fire, Burns, and Ineffective Energy Delivery

Do not discharge standard paddles on top of therapy electrodes or ECG electrodes. Do not allow standard paddles (or therapy electrodes) to touch each other, ECG electrodes, lead wires, dressings, transdermal patches, etc. Such contact can cause electrical arcing and patient skin burns during defibrillation and may divert defibrillating energy away from the heart muscle.

WARNING

Possible Skin Burns and Ineffective Energy Delivery

Therapy electrodes that are dried out or damaged may cause electrical arcing and patient skin burns during defibrillation. Do not use therapy electrodes that have been removed from foil package for more than 24 hours. Do not use electrodes beyond Use By date. Check that electrode adhesive is intact and undamaged. Replace adult therapy electrodes after 50 shocks or pediatric therapy electrodes after 25 shocks.

WARNING

Possible Skin Burns

During defibrillation or pacing, air pockets between the skin and therapy electrodes may cause patient skin burns. Apply therapy electrodes so that entire electrode adheres to skin. Do not reposition the electrodes once applied. If the position must be changed, remove and replace with new electrodes.

WARNING

Possible Skin Burns

Electrodes and cables that are not specified for use with the LIFEPAK 15 defibrillator may malfunction and cause skin burns. Use only the electrodes and cables that are specified for use with the LIFEPAK 15 defibrillator.

WARNING

Possible Defibrillator Shutdown

The large current draw required for defibrillator charging may cause the defibrillator to reach a shutdown voltage level with no low battery indication. If the defibrillator shuts down without warning or if a replace battery warning occurs, immediately replace the battery with another fully charged battery.

WARNING

Possible Interference with Implanted Electrical Device

Defibrillation may cause implanted devices to malfunction. Place standard paddles or therapy electrodes away from implanted devices if possible. Check implanted device function after defibrillation.

CAUTION

Possible Equipment Damage

Prior to using this defibrillator, disconnect from the patient all equipment that is not defibrillator-protected.

Therapy Electrode and Standard Paddle Placement

The following paragraphs describe therapy electrode and standard paddle skin preparation and placement, including special placement situations.

Patient Skin Preparation

Prepare the patient's skin:

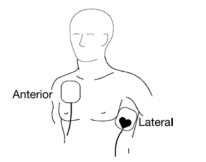
- Remove all clothing from the patient's chest.
- Remove excessive chest hair as much as possible. Avoid nicking or cutting the skin if using a shaver or razor. If possible, avoid placing electrodes over broken skin.
- Clean and dry the skin, if necessary. Remove any ointment on the patient's chest.
- Briskly wipe the skin dry with a towel or gauze. This mildly abrades the skin and removes oils, dirt, and other debris for better electrode adhesion to the skin.
- Do not use alcohol, tincture of benzoin, or antiperspirant to prep the skin.

Anterior-Lateral Placement

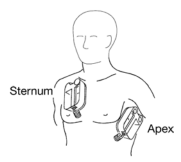
Anterior-lateral placement is used for ECG monitoring, defibrillation, synchronized cardioversion, and noninvasive pacing.

To perform anterior-lateral placement:

 Place either the ♥ therapy electrode or APEX paddle lateral to the patient's left nipple in the midaxillary line, with the center of the electrode in the midaxillary line, if possible. See the following figure.



QUIK-COMBO Therapy Electrodes



Standard Paddles

Figure 33 Anterior-Lateral Placement

2. Place the other therapy electrode or **STERNUM** paddle on the patient's upper right torso, lateral to the sternum and below the clavicle as shown in the preceding figure.

Anterior-Posterior Placement

Anterior-posterior is an alternative position for noninvasive pacing, manual defibrillation, and synchronized cardioversion, but not for ECG monitoring or AED mode. The ECG signal obtained through electrodes in this position is not a standard lead.

To perform anterior-posterior placement:

- 1. Place either the ♥ or + therapy electrode over the left precordium as shown in the following figure. The upper edge of the electrode should be below the nipple. Avoid placement over the nipple, the diaphragm, or the bony prominence of the sternum, if possible.
- 2. Place the other electrode behind the heart in the infrascapular area as shown in the following figure. For patient comfort, place the cable connection away from the spine. Do not place the electrode over the bony prominences of the spine or scapula.

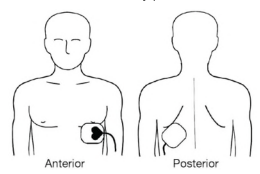


Figure 34 Anterior-Posterior Placement

Special Situations for Electrode or Paddle Placement

When placing therapy electrodes or standard paddles, be aware of the special requirements in the following possible situations.

Synchronized Cardioversion

Alternative placements for cardioversion of atrial fibrillation include a) place the ♥ therapy electrode over the left precordium and the other electrode on the patient's right posterior infrascapular area; or b) place the ♥ therapy electrode to the right of the sternum and the other electrode on the patient's posterior left infrascapular area.

Obese Patients or Patients with Large Breasts

Apply therapy electrodes or standard paddles to a flat area on the chest, if possible. If skin folds or breast tissue prevent good adhesion, it may be necessary to spread skin folds apart to create a flat surface.

Thin Patients

Follow the contour of the ribs and spaces when pressing therapy electrodes onto the torso. This action limits air spaces or gaps under the electrodes and promotes good skin contact.

Patients with Implanted Devices

Implanted devices such as cardiac defibrillators, pacemakers, or other devices may absorb energy from a LIFEPAK 15 defibrillator shock or be damaged by the shock. If possible, place therapy electrodes or standard paddles in the standard placements but away from the implanted device. Treat the patient like any other patient who requires care. If defibrillation is unsuccessful, it may be necessary to try alternate electrode placement (anterior-posterior).

Automated External Defibrillation (AED)

Intended Use

When used in AED mode, the LIFEPAK 15 monitor/defibrillator is a semiautomatic defibrillator that provides a prompted treatment protocol and ECG analysis using a patented Shock Advisory SystemTM (SAS). This software algorithm analyzes the patient's electrocardiographic (ECG) rhythm and indicates whether or not a shockable rhythm is detected. AED mode requires operator interaction in order to defibrillate the patient.

AED mode is intended for use by personnel who are authorized by a physician or medical director and have, at a minimum, the following skills and training:

- CPR training
- AED training equivalent to that recommended by the American Heart Association (AHA) or the European Resuscitation Council (ERC)
- Training in the use of the LIFEPAK 15 monitor/defibrillator in AED mode

Indications

AED mode is to be used only on patients in cardiopulmonary arrest. The patient must be unconscious, pulseless, and not breathing normally before using the defibrillator to analyze the patient's ECG rhythm. In AED mode, the LIFEPAK 15 monitor/defibrillator is not intended for use on pediatric patients less than eight years old.

Contraindications

None known.

AED Warnings

WARNING

Possible Misinterpretation of Data

Do not analyze in a moving vehicle. Motion artifact may affect the ECG signal resulting in an inappropriate **SHOCK** or **NO SHOCK ADVISED** message. Motion detection may delay analysis. Stop vehicle and stand clear of patient during analysis.

WARNING

Possible ECG Misinterpretation

Do not place therapy electrodes in the anterior-posterior position when operating this defibrillator in AED mode. A **SHOCK** or **NO SHOCK** decision may be inappropriately advised. The shock advisory algorithm requires the electrodes to be placed in the anterior-lateral (Lead II) position.

WARNING

Pediatric Patient Safety Risk

In AED mode, this defibrillator is not designed or tested to interpret pediatric rhythms or administer energy at pediatric joule settings for children under eight years old.

AED Mode

The LIFEPAK 15 monitor/defibrillator is set up to operate in Manual mode when it is turned on (factory default setting). The device can be set up to power on in AED mode by changing the Setup Options. The factory default settings for AED mode are identified in Setup Options Factory Default Settings (on page 239). The energy settings and other AED setup options can be changed according to medical protocol. For more information, see the LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device.

The ECG is continuously displayed in AED mode; however, access to other functions such as **OPTIONS** is not allowed in AED mode. The CPR metronome automatically sounds during CPR times, but it can only be silenced and un-silenced in AED mode. For more information, see CPR Time and Metronome (on page 125).

You can exit AED mode's prompted protocol and enter Advisory Monitoring or Manual Mode. For more information about Advisory Monitoring, see Advisory Monitoring (on page 129). Access to Manual mode may be direct, require confirmation or a passcode, or not allowed, depending on how your defibrillator is set up. It is important to be thoroughly familiar with your monitor/defibrillator settings and operation before use.

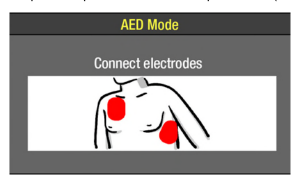
AED Procedure

The following descriptions of AED prompts (voice and text) are based on the factory default settings for AED mode. The settings are consistent with the 2010 American Heart Association (AHA) and European Resuscitation Council (ERC) guidelines. Changing the setup options may result in different AED behavior.

The CPR metronome automatically sounds during CPR times and can only be silenced and unsilenced.

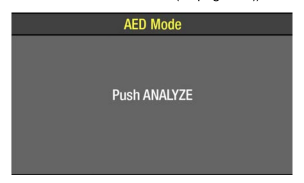
To perform automated external defibrillation:

- 1. Verify that the patient is in cardiopulmonary arrest (unconscious, pulseless, not breathing normally).
- 2. Press ON.
- 3. Prepare the patient for electrode placement (see Patient Skin Preparation (on page 117)).



The **CONNECT ELECTRODES** prompts occur until the patient is connected to the AED. If possible, place the patient on a hard surface away from standing water.

- 4. Connect the therapy electrodes to the therapy cable and confirm cable connection to the defibrillator.
- 5. Apply the therapy electrodes to the patient's chest in the anterior-lateral position (see Anterior-Lateral Placement (on page 117)).



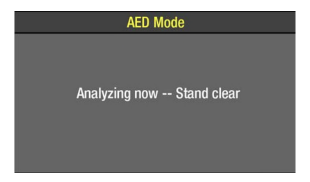
The **PUSH ANALYZE** prompts occur when the patient is properly connected to the AED.

6. Press ANALYZE to initiate the analysis. Stop CPR.

WARNING

Possible Misinterpretation of Data

Do not move the AED during analysis. Moving the AED during analysis may affect the ECG signal resulting in an inappropriate **SHOCK** or **NO SHOCK ADVISED** decision. Do not touch the patient or the AED during analysis.



The ANALYZING NOW—STAND CLEAR prompts occur. The SAS analyzes the patient's ECG in approximately 6 to 9 seconds and advises either SHOCK ADVISED or NO SHOCK ADVISED.

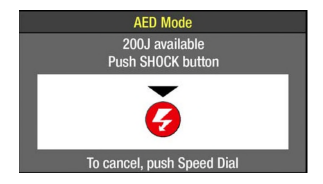
7. Continue to follow the screen messages and voice prompts provided by the AED.

Shock Advised

The following prompts occur when shock is advised:



If the AED detects a shockable rhythm, the **SHOCK ADVISED** prompts occur. Charging to the joule setting for Shock #1 begins. A charging bar appears and a ramping tone sounds.

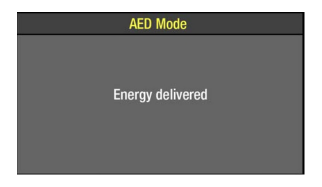


When charging is complete, the available energy is displayed.

The **STAND CLEAR, PUSH SHOCK BUTTON!** (►) message occurs, followed by a "Shock ready" tone.

Clear everyone away from touching the patient, bed, or any equipment that is connected to the patient.

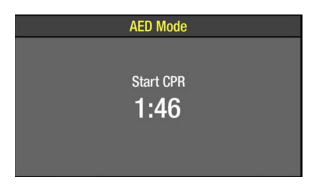
Press (shock) to deliver energy to the patient.



When the **ENERGY DELIVERED** message occurs indicating that the energy transfer was completed.

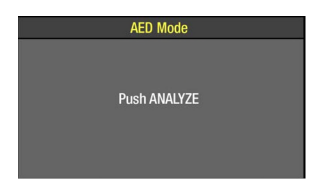


Note: If you do not press the (shock) button within 60 seconds, or the **SPEED DIAL** is pressed to cancel charging, the defibrillator disarms and the **DISARMING** message appears.



After a shock is delivered, the **START CPR** prompts occur. A countdown timer (min:sec format) continues for the duration specified in the **CPR TIME 1** setup option.

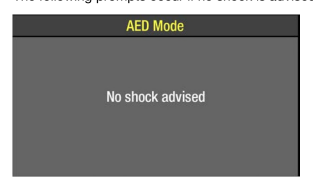
Note: The CPR metronome automatically provides audible compression "tocks" and ventilation prompts or tones only during CPR intervals at a ratio of 30:2. To silence the metronome, press **CPR**. To un-silence the metronome, press **CPR** again.



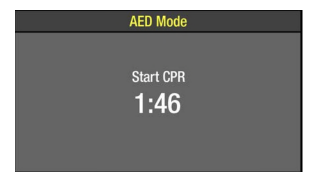
When the CPR countdown time ends, the **PUSH ANALYZE** prompts occur. These prompts repeat every 20 seconds until you press **ANALYZE**.

No Shock Advised

The following prompts occur if no shock is advised:

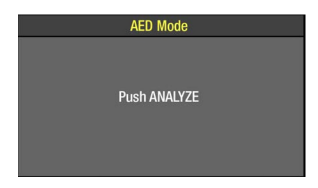


If the AED detects a nonshockable rhythm, the **NO SHOCK ADVISED** prompts occur. The defibrillator does not charge, and no shock can be delivered.



After NO SHOCK ADVISED, the START CPR prompts occur. A countdown timer (min:sec format) continues for the duration specified in the CPR TIME 2 setup option.

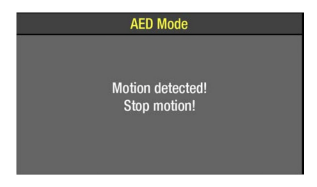
Note: The CPR metronome automatically provides audible compression "tocks" and ventilation prompts or tones only during CPR intervals. To silence the metronome, press **CPR**. To un-silence the metronome, press **CPR** again.



When the CPR countdown time ends, the **PUSH ANALYZE** prompts occur. These prompts repeat every 20 seconds until you press **ANALYZE**.

Subsequent analysis for **SHOCK ADVISED** and **NO SHOCK ADVISED** sequences are the same as described above. The energy level for Shock 2, 3, and greater depends on the **ENERGY PROTOCOL** setup and the analysis decision. When a **NO SHOCK ADVISED** decision follows a shock, the energy level does not increase for the next shock. When a **SHOCK ADVISED** decision follows a shock, the energy level increases for the next shock.

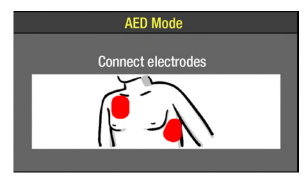
Motion Detected



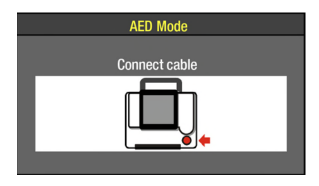
If the AED detects motion during the ECG analysis, the **MOTION DETECTED, STOP MOTION** prompts occur, followed by a warning tone.

Analysis is inhibited until the motion stops or for up to 10 seconds. After the motion ceases or 10 seconds have elapsed, analysis continues to completion even if motion is still present. For possible causes of motion detection and suggested solutions, see Troubleshooting Tips for AED Mode (on page 130).

Electrodes or Therapy Cable Off



If therapy electrodes are not connected, the **CONNECT ELECTRODES** prompts occur until the patient is connected.



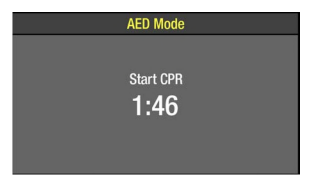
If the therapy cable is not connected to the defibrillator, the **CONNECT CABLE** message appears until the cable is connected.

Shock Counter



The shock counter (x) indicates how many shocks have been delivered to the patient. The shock counter resets to zero whenever the defibrillator is turned off for longer than 30 seconds.

CPR Time and Metronome



During use, CPR time shown on the countdown timer will vary slightly due to the metronome. When the CPR metronome is active during use, CPR times are adjusted to end CPR compression "tocks" on a compression cycle. As a result, the CPR countdown timer shows CPR times that approximate the seconds selected in Setup mode. Even if the metronome is off or silent during CPR time, the CPR time displayed will vary slightly from the time set up in Setup mode. This is because the metronome keeps track of compression "tocks" and ventilation prompts in the background so that if the metronome is activated, the CPR time ends with compressions.

Switching from AED Mode to Manual Mode

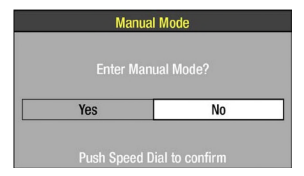
When in AED mode, Manual mode may be accessed directly, require confirmation or a passcode, or not be accessible at all depending on how your defibrillator has been set up.

To switch from AED mode to Manual mode, press **ENERGY SELECT** one time. You can also press **PACER** or **CHARGE** to switch from AED mode to Manual mode.

Note: If the metronome is active (providing compression "tocks" and ventilation prompts) when you switch from AED mode to Manual mode, the metronome stays active upon entering Manual mode.

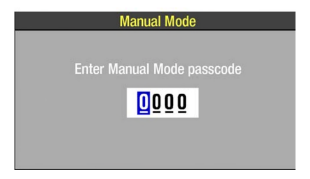
Depending on how manual access is set up, continue to Manual mode as follows:

- AED/Direct No restrictions to Manual mode access.
- **AED/Confirmed**—A confirmation screen appears:



Select YES to enter Manual mode.

• **AED/Passcode**—A passcode screen appears:



Rotate and press the **SPEED DIAL** to enter the passcode.

The code changes to dots to protect the passcode, and the defibrillator enters Manual mode.

You have three opportunities to enter the correct password. After an incorrect attempt, the message **INCORRECT--TRY AGAIN** appears. After three incorrect attempts, the message **ACCESS DENIED** appears, and the defibrillator returns to AED mode.

• **Restricted**—A **MANUAL MODE DISABLED** message appears, an alert tone sounds, and the LIFEPAK 15 monitor/defibrillator returns to AED mode.

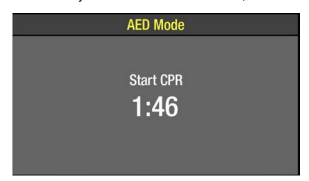
It is important that all users of the LIFEPAK 15 monitor/defibrillator be thoroughly familiar with the monitor/defibrillator settings and operation before use.

Special AED Setup Options

The following descriptions of AED prompts (voice and text) explain special setup options.

Initial CPR - CPR First

When the **INITIAL CPR** option is set to **CPR FIRST**, you are prompted to **START CPR** immediately after the AED is turned on, and before an analysis.



The **START CPR** prompts occur.



After 3 seconds, a countdown timer appears and the **IF YOU WITNESSED THE ARREST, PUSH ANALYZE** prompts occur. These prompts provide an opportunity to end the initial CPR early and proceed directly to analysis.

Note: The decision to end CPR early is based on your protocol and if you witnessed the arrest.

- If you did witness the arrest, press ANALYZE. The CPR period ends, and the ANALYZING NOW, STAND CLEAR prompts occur.
- If you did not witness the arrest, perform CPR and do not press ANALYZE. The Initial CPR
 countdown timer continues for the duration specified in the INITIAL CPR TIME setup
 option, for example, 90 seconds. When initial CPR time ends, the PUSH ANALYZE
 prompts occur.

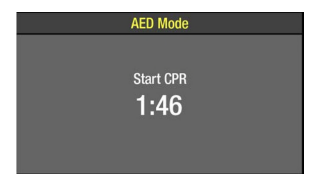
Initial CPR - Analyze First

When the **INITIAL CPR** option is set to **ANALYZE FIRST**, you are prompted to perform analysis after the AED is turned on. CPR is prompted after the AED completes the analysis.

If the electrodes are not attached to the patient, the **CONNECT ELECTRODES** prompts occur before you are prompted to perform analysis.

No Shock Advised If the AED detects a nonshockable rhythm, the **START CPR** prompts occur.

Automated External Defibrillation (AED)



A countdown timer (min:sec format) continues for the duration specified in the **INITIAL CPR TIME** setup option.

When initial CPR time ends, the **NO SHOCK ADVISED** prompts occur, followed by **PUSH ANALYZE**.

Shock Advised If the AED detects a shockable rhythm, the **START CPR** prompts occur, followed by **IF YOU WITNESSED THE ARREST, PUSH ANALYZE**.



These prompts provide an opportunity to end the initial CPR early and proceed directly to delivering a shock.

Note: The decision to end CPR early is based on your protocol and if you witnessed the arrest.

- If you did witness the arrest, press ANALYZE. This ends the initial CPR period and the SHOCK ADVISED and STAND CLEAR, PUSH SHOCK button! (*) prompts occur. Proceed according to your training with the AED for delivering the shock.
- If you did not witness the arrest, perform CPR and do not press ANALYZE to end CPR early. The Initial CPR countdown timer continues for the duration specified in the INITIAL CPR TIME setup option, for example, 90 seconds. Near the end of CPR time, the defibrillator silently charges to prepare for the shock. CPR continues up to shock delivery. When initial CPR time ends, the SHOCK ADVISED and STAND CLEAR, PUSH SHOCK button! () prompts occur. Proceed according to your training with the AED for delivering a shock.

Pre-shock CPR Time

When **PRE-SHOCK CPR** time is set to 15 seconds or more, you are prompted to start CPR immediately after a shockable rhythm is detected, before the shock is delivered.



After analysis is complete, the **START CPR** prompts occur. A countdown timer (min:sec format) continues for the duration specified in the **PRE-SHOCK CPR** time setup option.

The defibrillator silently charges in preparation for the shock.

When CPR time ends, the SHOCK ADVISED and STAND CLEAR, PUSH

SHOCK BUTTON! (₹) prompts occur. Proceed according to your training with the AED for delivering a shock.

Note: The **E** (shock) button is disabled during the pre-shock CPR interval to avoid accidental shock delivery while the defibrillator is charged and a responder is performing CPR.

Advisory Monitoring

Advisory Monitoring is a special way to set up AED mode that allows the use of all the monitoring functions without initiating the AED prompted protocol when the device is turned on. When needed, the AED mode prompted protocol can be initiated by pressing **ANALYZE**. In addition, access to Manual mode therapies—that is, manual defibrillation, synchronized cardioversion, or pacing—by unauthorized users can be restricted, if necessary.

Certain setup options must be changed for the device to operate in Advisory Monitoring when it is turned on. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

When set up for Advisory Monitoring and the monitor is turned on, the **ADVISORY MODE-MONITORING** message appears continuously in the message area on the Home Screen. Monitor functions such as NIBP, SpO₂ and 12-lead ECG can be used. Lead II and dashes are shown in the top ECG trace (Channel 1) unless or until the patient is connected to the ECG cable. If therapy electrodes (pads) and the therapy cable are connected to the patient, press **LEAD** to change to **PADDLES** lead and view the ECG.

In Advisory Monitoring, **LEAD II** and **PADDLES** lead are the only ECG monitoring leads allowed in Channel 1. The Continuous Patient Surveillance System (CPSS) is active and automatically evaluates the patient ECG. However, CPSS is evaluating only for a potentially shockable rhythm. If a shockable ECG rhythm such as VF is detected, the following prompt appears: **CHECK PATIENT. IF NO PULSE, PUSH ANALYZE**.

Prior to pressing **ANALYZE**, confirm that the patient is in cardiac arrest. Motion artifact, a low amplitude ECG, and other causes of poor ECG signal may cause false CPSS alerts. If the patient is not in cardiac arrest, do not press **ANALYZE**. Troubleshoot the cause of the false CPSS alert.

If the patient is in cardiac arrest, press **ANALYZE**. Pressing **ANALYZE** causes the defibrillator to enter AED mode. The defibrillator begins the AED prompted protocol and analyzes the

Automated External Defibrillation (AED)

patient's ECG when therapy electrodes are applied to the patient. For more information about defibrillator behavior in AED mode, see Automated External Defibrillation (AED) (on page 119).

Note: CPSS only evaluates for shockable ECG rhythms. If the ECG rhythm is nonshockable, for example asystole, no prompting occurs. Users who are not trained to interpret ECGs or are trained only to use AED mode must always press ANALYZE when using this special setup function to initiate ECG analysis and AED prompting.

To switch back to Advisory Monitoring from AED prompted protocol, press **LEAD**.

For information about limiting access to Manual mode by unauthorized users, see Switching from AED Mode to Manual Mode (on page 126), or see the LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device.

Troubleshooting Tips

| Table 21 Troubleshooting Tips for AED Mode | | | |
|---|---|---|--|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION | |
| message appears | Therapy electrodes are not connected to the therapy cable | Check for electrode connection. | |
| | Electrodes do not adhere properly to the patient | Press electrodes firmly on patient's skin. Clean, shave, and dry the patient's skin as recommended. Replace the electrodes. | |
| | Electrodes are dry, damaged, or out of date | Apply new electrodes. | |
| | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. | |
| CONNECT CABLE message appears | Therapy cable is disconnected during charging | Reconnect cable and press CHARGE again. | |
| | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. | |
| MOTION DETECTED and STOP MOTION messages appear during analysis | Patient movement | Stop CPR during analysis. When patient is being manually ventilated, press ANALYZE after complete exhalation. | |
| | Patient movement because of agonal respirations | Allow analysis to proceed to completion—analysis is delayed no more than 10 seconds due to motion detection. | |

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|--|---|
| VECENTION | Electrical/radio frequency interference | Move hand-held communication devices or other suspected devices away from the defibrillator, when possible. |
| | Vehicle motion | Stop vehicle during analysis.Move patient to stable location, when possible. |
| DISARMING message appears (energy charge removed) | (shock) button not pressed within 60 seconds after charge complete | Recharge the defibrillator, if desired. |
| | SPEED DIAL pressed | Recharge the defibrillator. |
| | Therapy electrodes or cable disconnected | Reconnect electrode or cable. |
| Energy did not escalate | After a shock, the next analysis was NO SHOCK ADVISED | No action needed. Defibrillator does not escalate energy when a NO SHOCK ADVISED decision follows a shock. |
| Charge time to 360 joules exceeds 10 seconds | Battery low | Replace battery with fully charged battery. Connect to auxiliary power using approved power adapter. |
| | Operating temperature is too low | Move patient and device to warmer environment, if necessary. |
| REPLACE BATTERY prompt occurs | Both batteries are very low | Replace one or both batteries immediately. Connect to auxiliary power using approved power adapter. |
| Voice prompts sound faint or distorted | Low battery power | Replace one or both batteries immediately. Connect to auxiliary power using approved power adapter. |
| CPR time shown (minutes/seconds) is different than expected | Function of metronome | None. The metronome adjusts the CPR time to ensure CPR cycle ends with compressions. (See CPR Time and Metronome (on page 125).) |
| | Incorrect setup option selected | Change CPR time setup option. See LIFEPAK 15 Monitor/Defibrilla tor Setup Options provided with your device. |
| Press CPR and metronome does not activate | In AED mode, and not in CPR interval | Wait until CPR interval (audible "tocks") to silence or activate metronome. |

Manual Defibrillation

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|---|---|
| Home Screen is blank but ON LED is illuminated | Screen not functioning properly | Press ANALYZE and follow voice prompts to treat patient. |
| Analysis result is NO SHOCK ADVISED and ECG shows a perfectly flat, isoelectric line. | The Test Load is connected to therapy cable | Remove the Test Load and connect therapy electrodes to the cable. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Manual Defibrillation

The LIFEPAK 15 monitor/defibrillator provides manual defibrillation using adult and pediatric QUIK-COMBO pacing/defibrillation/ECG electrodes, adult standard paddles, or pediatric paddles. For more information, see Paddle Accessory Options (on page 149).

The LIFEPAK 15 monitor/defibrillator is capable of providing intra-operative direct defibrillation and synchronized cardioversion with the internal paddles accessory designed for the LIFEPAK 15 defibrillator. For more information, see the Instructions for Use for the internal paddles.

Intended Use

When used in Manual mode, the LIFEPAK 15 monitor/defibrillator is a direct current defibrillator that applies a brief, intense pulse of electricity to the heart muscle. Manual mode requires operator interpretation of the ECG rhythm and interaction with the device in order to defibrillate the patient.

Manual mode defibrillation and synchronized cardioversion are intended for use by personnel who are authorized by a physician or medical director and have, at a minimum, the following skills and training:

- Arrhythmia recognition and treatment
- · Advanced resuscitation training equivalent to that recommended by the AHA or ERC
- Training on the use of the LIFEPAK 15 monitor/defibrillator

Defibrillation is only one aspect of the medical care required to resuscitate a patient who has a shockable ECG rhythm. Depending on the situation, other supportive measures may include:

- Cardiopulmonary resuscitation (CPR)
- Administration of supplemental oxygen
- Drug therapy

Indications

Manual defibrillation is indicated for the termination of certain potentially fatal arrhythmias, such as ventricular fibrillation and symptomatic ventricular tachycardia. Delivery of this energy in the synchronized mode is a method for treating atrial fibrillation, atrial flutter,

paroxysmal supraventricular tachycardia and, in relatively stable patients, ventricular tachycardia.

Contraindications

Defibrillation is contraindicated in the treatment of Pulseless Electrical Activity (PEA), such as idioventricular or ventricular escape rhythms, and in the treatment of asystole.

Manual Defibrillation Warnings

WARNING

Shock Hazard

Conductive gel (wet or dry) on the paddle handles can allow the electrical energy to discharge through the operator during defibrillation. Completely clean the paddle electrode surfaces, handles, and storage area after defibrillation.

WARNING

Possible Fire, Burns, and Ineffective Energy Delivery

Precordial lead electrodes and lead wires may interfere with the placement of standard paddles or therapy electrodes. Before defibrillation, remove any interfering precordial lead electrodes and lead wires.

WARNING

Possible Burns and Ineffective Energy Delivery

A gel pathway on the skin between the standard paddles will cause defibrillating energy to arc between paddles and divert energy away from the heart muscle. Do not allow conductive gel (wet or dry) to become continuous between paddle sites.

WARNING

Possible Patient Skin Burns

During defibrillation, air pockets between the skin and standard paddles can cause patient skin burns. Completely cover paddle electrode surfaces with fresh conductive gel and apply 25 lb of pressure per paddle during discharge.

WARNING

Possible Paddle Damage and Patient Skin Burns

Discharging the defibrillator with the standard paddle surfaces shorted together can pit or damage the paddle electrode surface. Pitted or damaged paddle surfaces may cause patient skin burns during defibrillation. Discharge the defibrillator only as described in these operating instructions.

WARNING

Possible Incorrect Energy Delivery

The defibrillator does not automatically adjust energy when using pediatric therapy electrodes or pediatric standard paddles. Manually select the appropriate energy prior to defibrillating the patient.

Manual Mode

The LIFEPAK 15 monitor/defibrillator is set up to operate in Manual mode when it is turned on (factory default setting). If required by your protocols, the defibrillator can be set up to power on in the automated external defibrillator (AED) mode. For information on switching from AED mode to Manual mode, see Switching from AED Mode to Manual Mode (on page 126).

Manual Defibrillation Procedure

To perform manual defibrillation:

- 1. Verify that the patient is in cardiopulmonary arrest (unconscious, pulseless, not breathing normally).
- 2. Press ON.
- Identify the electrode or paddle sites on the patient and prepare the patient's skin. (See Patient Skin Preparation (on page 117).) Use either the anterior-lateral or anterior-posterior position.
- 4. Connect the therapy electrodes to the therapy cable and confirm cable connection to the defibrillator.
- 5. Apply therapy electrodes to the patient in anterior-lateral or anterior-posterior position. If using standard paddles, apply conductive gel to the paddles and place paddles on the patient's chest in the anterior-lateral position.
- 6. Confirm desired energy is selected, or press **ENERGY SELECT** or rotate the **SPEED DIAL** to select the desired energy. On the standard (hard) paddles, rotate the **ENERGY SELECT** dial.
- 7. Press **CHARGE**. While the defibrillator is charging, a charging bar appears and a ramping tone sounds, indicating the charging energy level. When the defibrillator is fully charged, the screen displays available energy.
- 8. Make certain all personnel, including the operator, stand clear of the patient, stretcher, bed, and any equipment connected to the patient.
- 9. Confirm ECG rhythm requires defibrillation. Confirm available energy.

10. Press the

(shock) button on the defibrillator or the

(shock) buttons on the standard paddles to discharge energy to the patient. For standard paddles, apply firm pressure with both paddles to the patient's chest, and press both paddle buttons simultaneously to discharge energy to the patient. For safety reasons, the

(shock) button on the defibrillator front panel is disabled when using standard paddles.

Note: To disarm (cancel the charge), press the **SPEED DIAL**. The defibrillator disarms automatically if shock buttons are not pressed within 60 seconds, or if you change the energy selection after charging begins.

Note: To interrupt defibrillation and initiate pacing, press **PACER**. If charged, the defibrillator disarms.

- 11. Start CPR according to your protocol. To activate the metronome, press CPR at any time.
- 12. At the end of your CPR period, observe the patient and the ECG rhythm. If an additional shock is necessary, repeat the procedure beginning at Step 6.

Successful resuscitation is related to the length of time between the onset of a heart rhythm that does not circulate blood (ventricular fibrillation, pulseless ventricular tachycardia) and defibrillation. The physiological state of the patient may affect the likelihood of successful defibrillation. Thus, failure to resuscitate a patient is not a reliable indicator of defibrillator performance. Patients often exhibit a muscular response (such as jumping or twitching) during an energy transfer. The absence of such a response is not a reliable indicator of actual energy delivery or device performance.

Using the CPR Metronome

When CPR is required during cardiac arrest, the CPR metronome provides audible prompts that guide the user to deliver CPR with proper timing in accordance with the 2010 American Heart Association and European Resuscitation Council CPR guidelines.

CPR Metronome Warnings

WARNING

CPR Delivered When Not Needed

The metronome sounds do not indicate information regarding the patient's condition. Because patient status can change in a short time, the patient should be assessed at all times. Do not perform CPR on a patient who is responsive or is breathing normally.

Note: The CPR metronome is a tool to be used as a timing aid during CPR. Assess the patient at all times and provide CPR only when indicated. Provide CPR according to your training and protocols.

How the CPR Metronome Works

The metronome provides audible "tocks" at a rate of 100/minute to guide the rescuer in performing chest compressions. The metronome also provides audible ventilation prompts (either a tone or verbal "ventilate") to cue the rescuer when to provide ventilations. The

metronome prompts the rescuer to perform CPR at the selected compression to ventilation (C:V) ratio.

Age-Airway Considerations

The default C:V ratio for the metronome (in both AED and Manual modes) is Adult - No Airway (30:2) because most patients in cardiac arrest are adults who have an initially unsecured airway. In Manual mode, the user can choose the most appropriate C:V ratio based on the patient's age and current airway status. The Age-Airway selection determines the C:V ratio of the metronome sounds. The default C:V ratios are shown in the following table.

Table 22 Default Age-Airway C:V Ratios in Manual Mode

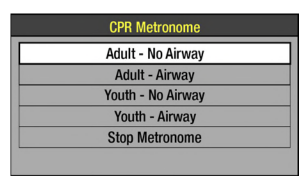
| AGE-AIRWAY | C:V RATIO |
|----------------------|-----------|
| Adult - No Airway* | 30:2 |
| Adult - Airway** | 10:1 |
| Youth - No Airway*** | 15:2 |
| Youth - Airway | 10:1 |

^{*} No Airway = No artificial airway in place

Note: The compression-to-ventilation ratio selections can be set up according to local medical protocols. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

Activating and Deactivating the Metronome

To activate the CPR metronome in Manual mode:



- Press CPR. The CPR Metronome menu appears and the metronome is activated using the Adult-No Airway default setting.
- 2. Use the **SPEED DIAL** to highlight and select the desired Age-Airway setting.

CPR: Adult - No Airway 30:2

When the metronome is on, a message appears in the message area that indicates the current Age-Airway selection.

Note: If the VF/VT alarm is on, it is suspended when the metronome is on to prevent false VF/VT alarms. If other vital sign alarms activate when the metronome is on, the visual indicators occur, but the alarm tone is suppressed until the metronome is deactivated.

^{**} Airway = Advanced artificial airway in place

^{***} Youth = Pre-pubescent child

The metronome provides "tocks" and ventilation prompts continuously until it is deactivated. To stop the metronome, select **STOP METRONOME** in the CPR Metronome menu. An event is recorded in the CODE SUMMARY Event Log when CPR metronome is turned ON or OFF and when the Age-Airway setting is changed. To adjust the volume of the metronome, press **OPTIONS**, select **ALARM VOLUME**, and change the **VOLUME**.

Note: If all Age-Airway selections are set to the same C:V ratio (for example, Adult - No Airway, Adult - Airway, Youth - No Airway, and Youth - Airway all set to 10:1), the CPR metronome always provides "tocks" and ventilation prompts at the set ratio for both AED mode and Manual mode. In this situation, the CPR Metronome menu does not appear when **CPR** is pressed during use—pressing the **CPR** button only activates and deactivates the metronome at the fixed C:V ratio.

Synchronized Cardioversion Procedure

The LIFEPAK 15 monitor/defibrillator can be set up to remain in Sync mode or to return to Asynchronous mode after a shock is delivered. The factory default setting is to return to Asynchronous mode after a shock. It is important that you know how your defibrillator is set up. For information about changing the setup option, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

To perform synchronized cardioversion:

- 1. Press ON.
- 2. Attach patient ECG cable and ECG electrodes as previously described (see Monitoring the ECG (on page 47)). ECG electrodes and cable must be used to monitor the ECG when standard paddles are used for cardioversion.
- 3. Select Lead II or lead with greatest QRS complex amplitude (positive or negative).

Note: To monitor the ECG using therapy electrodes, place the electrodes in anterior-lateral position and select **PADDLES** lead.

WARNING

Possible Lethal Arrhythmia

Ventricular fibrillation may be induced with improper synchronization. DO NOT use the ECG from another monitor (slaving) to synchronize the monitor/defibrillator's discharge. Always monitor the patient's ECG directly through the defibrillator's ECG cable or therapy cable. Confirm proper placement of the sense markers on the ECG.

4. Press SYNC. The SYNC MODE message appears in the message area when Sync is active.

Note: Press **SYNC** again to deactivate Sync mode.

Synchronized Cardioversion Procedure

- 5. Observe the ECG rhythm. Confirm that a triangle sense marker (▼) appears near the middle of each QRS complex. If the sense markers do not appear or are displayed in the wrong locations (for example, on the T-wave), adjust ECG SIZE or select another lead. (It is normal for the sense marker location to vary slightly on each QRS complex.)
- Connect the therapy electrodes to the therapy cable and confirm cable connection to the defibrillator.
- 7. Prepare the patient's skin and apply therapy electrodes to the patient in the anterior-lateral position. (See Therapy Electrode and Standard Paddle Placement (on page 116).) If using standard paddles, apply conductive gel to the paddles and place paddles on the patient's chest.
- 8. Press **ENERGY SELECT** or rotate the **SPEED DIAL** to select the desired energy. On the standard (hard) paddles, rotate the **ENERGY SELECT** dial.
- Press CHARGE. While the defibrillator is charging, a charging bar appears and a ramping tone sounds, indicating the charging energy level. When the defibrillator is fully charged, the screen displays available energy.
- 10. Make certain all personnel, including the operator, stand clear of the patient, bed, stretcher, and any equipment connected to the patient.
- 11. Confirm ECG rhythm. Confirm available energy.
- 12. Press and *hold* the

 (shock) button on the defibrillator until the **ENERGY DELIVERED**message appears on the screen. For standard paddles, press and hold both

 (shock) buttons on the paddles simultaneously until the **ENERGY DELIVERED** message appears on the screen. Release buttons. For safety reasons, the

 (shock) button on the defibrillator front panel is disabled when using standard paddles.

Note: To disarm (cancel a charge), press the **SPEED DIAL**. The defibrillator disarms automatically if shock buttons are not pressed within 60 seconds, or if you change the energy selection after charging begins.

13. Observe patient and ECG rhythm. Repeat procedure starting from Step 4, if necessary.

Troubleshooting Tips

Table 23 Troubleshooting Tips for Defibrillation and Synchronized Cardioversion

| OBSERVATION | r Defibrillation and Synchronized C POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|---|--|
| Charge time to 360 joules exceeds 10 seconds | Battery low | Replace battery with fully charged battery. |
| | Operating temperature is too low | Move patient and device to warmer environment, if necessary. |
| Energy not delivered to patient when (shock) buttons are pressed | Device is in Sync mode and QRS complexes are not detected | Adjust ECG size for optimum sensing QRS or deactivate SYNC if rhythm VF/VT. |
| | SYNC accidentally pressed and rhythm is VF/VT | Press SYNC to turn off Sync.Press (shock) buttons. |
| | Device in Sync mode and (shock) buttons not pressed and held until next detected QRS | Hold (shock) buttons until discharge occurs or next detected QRS and ENERGY DELIVERED message appears. |
| | (shock) buttons pressed before full charge reached | Wait for tone and message indicating full charge. |
| | Standard paddles connected and (shock) button on defibrillator front panel pressed | Simultaneously press (shock) buttons on standard paddles to discharge. |
| | Sixty seconds elapsed before (shock) buttons were pressed after full charge. Energy was internally removed. | Press (shock) buttons within 60 seconds of full charge. |
| | Energy selection changed | • Press CHARGE again. |
| CONNECT CABLE message appears | Therapy cable disconnected during charging | Reconnect cable and press CHARGE again. |
| | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. |
| ENERGY FAULT message appears (selected and available energy) | Defibrillator out of calibration | Attempt to transfer energy.Contact a qualified service technician. |
| DISARMING message appears | (shock) button not pressed within 60 seconds after charge complete | Recharge the defibrillator, if desired. |
| | Energy selected after charge complete | Recharge the defibrillator. |
| | SPEED DIAL pressed | Recharge the defibrillator. |
| | PACER pressed | Recharge, if necessary, or no action, if pacing desired. |
| | Therapy electrodes or cable disconnected | Reconnect electrode or cable. |

Synchronized Cardioversion Procedure

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|--|---|---|
| Energy did not escalate automatically per energy protocol | ENERGY SELECT pressed and disabled automatic protocol | Continue to select energy manually to treat patient. For more information about energy protocol, see LIFEPAK 15 Monitor/Defibrill ator Setup Options provided with your device. |
| SYNC mode will not activate | PACER is on. Pacing and Sync are separate functions and are not allowed at the same time. | Discontinue pacing, if appropriate for the patient, and press SYNC. |
| | ECG electrodes not attached to patient and standard paddles connected to defibrillator | Connect ECG electrodes to patient. |
| Patient did not "jump" (no muscle response) during defibrillator discharge | Patient muscle response is variable and depends on patient condition. Lack of visible response to defibrillation does not necessarily mean the discharge did not occur. | No action needed. |
| | The Test Load is connected to therapy cable | Remove the Test Load and connect therapy electrodes to cable. |
| ABNORMAL ENERGY DELIVERY message appears and Shock XJ Abnormal annotated on printout | Open air discharge with standard paddles | Press paddles firmly on patient's chest when discharging. |
| | Standard paddles placed face- to-face when (shock) button pressed | Perform test discharges per Operator's Checklist. See Manual Defibrillation Warnings (on page 133). |
| | Patient impedance is out of range | Increase energy or repeat shocks as needed. Consider replacing disposable therapy electrodes with new ones. |
| | Internal fault occurred | Repeat shock. Perform CPR and obtain another defibrillator, if necessary. |
| CONNECT ELECTRODES message appears | Therapy electrodes are not connected to the therapy cable | Check for electrode connection. |
| | Electrodes do not adhere properly to the patient | Press electrodes firmly on patient's skin. Clean, shave, and dry the patient's skin as recommended. Apply new electrodes. |
| | Electrodes are dry, damaged, or out of date | Apply new electrodes. |

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Chapter 5 | Therapy

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|---------------------------------|--|
| | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. |
| REPLACE BATTERY prompt occurs | Both batteries are very low | Replace one or both batteries immediately. |
| | | Connect to auxiliary power using approved power adapter. |
| CPR time shown (minutes/seconds) is different than expected | Metronome is on | None. The metronome adjusts the CPR time to ensure CPR cycle ends with compressions. |
| | Incorrect setup option selected | Change CPR time setup option. See LIFEPAK 15 Monitor/Defibrill ator Setup Options provided with your device. |
| Home Screen is blank but ON LED is illuminated | Screen not functioning properly | Print ECG strip to assess rhythm and other active vital signs. |
| | | Press ANALYZE and use AED mode, if necessary. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Noninvasive Pacing

The LIFEPAK 15 monitor/defibrillator provides noninvasive pacing using adult or pediatric QUIK-COMBO pacing/defibrillation/ECG electrodes. For more information, see Paddle Accessory Options (on page 149).

Intended Use

A noninvasive pacemaker is a device that delivers an electrical stimulus to the heart causing cardiac depolarization and myocardial contraction. The energy is delivered through large adhesive electrodes placed on the chest. In addition to noninvasive pacing, other supportive measures may be necessary.

Noninvasive pacing is intended for use by personnel who are authorized by a physician or medical director and have, at a minimum, the following skills and training:

- Arrhythmia recognition and treatment
- Advanced resuscitation training equivalent to that recommended by the AHA or ERC
- Training on the use of the LIFEPAK 15 monitor/defibrillator

Indications

Noninvasive pacing is indicated for symptomatic bradycardia in patients with a pulse.

Contraindications

Noninvasive pacing is contraindicated for the treatment of ventricular fibrillation and asystole.

Noninvasive Pacing Warnings

WARNING

Possible Inability to Pace

Using other manufacturers' combination therapy electrodes with this device could cause a decrease in pacing efficacy or the inability to pace because of unacceptably high impedance levels and invalidate the safety agency certifications. Use only the therapy electrodes that are specified in these operating instructions.

Demand and Nondemand Pacing

The LIFEPAK 15 monitor/defibrillator can be used for either demand or nondemand (asynchronous or "fixed rate") pacing.

Demand mode is used for most patients. In demand mode, the LIFEPAK 15 pacemaker inhibits pacing output when it "senses" the patient's own beats (intrinsic QRSs). In demand mode, if the

ECG SIZE is set too low to detect the patient's beats, or if an ECG lead becomes detached so that the ECG rhythm is not present, the pacemaker generates pacing pulses asynchronously. This means that the pacemaker generates pacing pulses at the selected rate regardless of the patient's ECG rhythm.

Nondemand mode can be selected if noise or artifact interferes with proper sensing of QRS complexes. Press **OPTIONS** to access nondemand mode. For more information, see Options (on page 41).

Noninvasive Pacing Procedure

ECG monitoring during pacing is performed with the ECG electrodes and patient ECG cable. Therapy electrodes are not capable of monitoring ECG and delivering pacing current at the same time.

Be sure to place the QUIK-COMBO therapy electrodes in the proper locations. Improper placement of the electrodes may make a difference in the capture threshold. For example, if the electrode placement is reversed, more pacing current may be needed to achieve capture.

WARNING

Possible Interruption of Therapy

Observe the patient continuously while the pacemaker is in use. Patient response to pacing therapy (for example, capture threshold) may change over time.

To perform noninvasive pacing:

- 1. Press ON.
- 2. Connect the patient ECG cable, apply ECG electrodes to the ECG cable and patient, and select Lead I, II, or III. To receive the best monitoring signal, make sure there is adequate space between the ECG electrodes and the therapy electrodes.
- 3. Identify the QUIK-COMBO therapy electrode sites on the patient. Use either the anterior-lateral or anterior-posterior position and prepare the patient's skin. (See Therapy Electrode and Standard Paddle Placement (on page 116).)
- 4. Apply therapy electrodes to the patient.
- 5. Connect the therapy electrodes to the therapy cable.
- 6. Press PACER.

WARNING

Possible Ineffective Pacing

The ECG size must be properly adjusted so that the patient's own beats are detected. If ECG size is set too high or too low, pacing pulses may not be delivered when required. Adjust ECG size so that sense markers are placed on the patient's QRS complexes.

Noninvasive Pacing

- 7. Observe the ECG rhythm. Confirm that a triangle sense marker () appears near the middle of each QRS complex. If the sense markers do not appear or are displayed in the wrong location (for example, on the T-wave), adjust ECG SIZE, or select another lead. (The sense marker location may vary slightly on each QRS complex.)
- 8. Press RATE or rotate the SPEED DIAL to select the desired pacing rate.
- Press CURRENT or rotate the SPEED DIAL to increase current until electrical capture occurs.
 Electrical capture is indicated by a wide QRS complex and a T-wave following the pace
 marker. For each delivered pacing stimulus, a positive pace marker displays on the ECG
 waveform.

Note: Dashes (---), not heart rate, are displayed on the Home Screen during noninvasive pacing, and heart rate alarms are disabled.

10. Palpate patient's pulse or check blood pressure to assess for mechanical capture. Consider use of sedation or analgesia if patient is uncomfortable.

Note: To change rate or current during pacing, press **RATE** or **CURRENT**. The **RATE** and **CURRENT** buttons allow changes in increments of 10; the **SPEED DIAL** allows changes in increments of 5.

Note: To interrupt pacing and view the patient's intrinsic rhythm, press and hold **PAUSE**. This causes the pacer to pace at 25% of the set rate. Release **PAUSE** to resume pacing at the set rate.

11. To stop pacing, reduce current to zero or press **PACER**.

Note: To defibrillate and stop noninvasive pacing, press **CHARGE**. Pacing automatically stops. Proceed with defibrillation.

The physiologic state of the patient may affect the likelihood of successful pacing or of skeletal muscle activity. The failure to successfully pace a patient is not a reliable indicator of pacemaker performance. Similarly, the patient's muscular response to pacing is not a reliable indicator of current delivered.

WARNING

Possible Patient Skin Burns

Prolonged noninvasive pacing may cause patient skin irritation and burns, especially with higher pacing current levels. Discontinue noninvasive pacing if skin becomes burned and another method of pacing is available. For additional information about therapy electrodes, see QUIK-COMBO Therapy Electrodes (on page 151).

If the monitor detects **ECG LEADS OFF** during pacing, pacing automatically switches to nondemand and continues at a fixed rate until the ECG lead is reattached. During nondemand pacing, the pacemaker delivers pulses at the set pace rate regardless of any intrinsic beats that the patient may have. The monitor continues to display the pacing rate (ppm) and the current (mA). To reestablish demand pacing, reattach the ECG lead.

While pacing, visually monitor the patient at all times—do not rely on the **ECG LEADS OFF** warning to detect changes in pacing function. Routinely assess for proper ECG sensing, pace pulse delivery, electrical capture, and mechanical capture.

If pacing electrodes detach during pacing, you see **CONNECT ELECTRODES** and **PACING STOPPED** messages and hear an alarm. The pacing rate is maintained and the current resets to 0 mA. Reattaching the pacing electrodes silences the alarm and removes the **CONNECT ELECTRODES** message. The current remains at 0 mA until you increase the current manually.

To turn off the LIFEPAK 15 monitor/defibrillator, pacing must be stopped. If the **ON** button is pressed when **PACER** is active, an alert tone sounds and the **PACING IN PROGRESS** message appears.

Troubleshooting Tips

Table 24 Troubleshooting Tips for Noninvasive Pacing

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|--|--|
| Device does not function when PACER is pressed | Power off | Check if power is ON . |
| | Low battery | Replace battery with fully charged battery. |
| PACER LED is on, but | Therapy electrodes off | Check for message displayed. |
| CURRENT (mA) will not increase | | Inspect therapy cable and electrode connections. |
| PACER LED on, CURRENT (mA) >0, but pace markers absent (not pacing) | Pacing rate set below patient's intrinsic rate | • Increase PPM. |
| | Pacer oversensing (ECG artifact, ECG size too high) | Establish clean ECG; decrease ECG size. |
| | | Select nondemand pacing. |
| Monitor screen displays distortion while pacing | ECG electrodes not optimally placed with respect to pacing | Reposition electrodes away from pacing electrodes. |
| | electrodes | • Select another lead (I, II, or III). |
| Pacing stops spontaneously | PACER pressed off | Press PACER and increase the current. |
| | Internal error detected. Service | Check for service indicator. |
| | message indicates an internal failure. | Cycle power and start pacing again. |
| | | Obtain service by a qualified service technician. |
| | Therapy electrode off | Check for message. Check pacing cable and electrode connections. |
| | CHARGE pressed | Press PACER and increase current, if pacing desired. Otherwise, proceed with defibrillation. |
| | Radio frequency interference | Move radio equipment away from pacemaker. |

Noninvasive Pacing

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|---|--|
| No muscle response to pacing | Patient's heart rate may be greater than noninvasive pacer ppm | No action needed. |
| | The Test Load is connected to therapy cable | Remove the Test Load and connect therapy electrodes to cable. |
| | Patient muscle response is variable and depends on patient condition. Muscular response to pacing is not a reliable indicator of current delivered. | No action needed. |
| Capture does not occur with pacing stimulus | Current (mA) set too low | Increase pacing current. (Administer sedation or analgesia as needed.) |
| CONNECT CABLE or PACING STOPPED message appears | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. |
| CONNECT ELECTRODES message appears | Pacing cable or electrode disconnected | Reconnect and set current. |
| | Electrodes not adhering to skin | Prepare skin. |
| | Therapy cable damaged | Replace therapy cable and perform daily checks per Operator's Checklist. |
| | Electrodes outdated | Replace electrodes and set current. |
| PACING IN PROGRESS message appears | CPR pressed | Press PACER to stop pacing, if appropriate, and then press CPR. |
| Pacing stops spontaneously and PACER FAULT message appears | Internal error detected | Cycle power and start pacing again.Obtain service by a qualified |
| | | service technician. |
| Intrinsic QRS complexes not sensed when pacing | ECG size too low | Increase ECG size or select another lead. |
| | Intrinsic QRS complexes are occurring during pacemaker's refractory period | Adjust PPM. |
| Pacing starts spontaneously | Patient's heart rate falls below set pacing rate | Appropriate pacemaker function; assess patient. |
| | During standby pacing, ECG lead disconnects and pacing begins asynchronously | Reconnect ECG lead. |
| Set pacing rate (ppm) and ECG paced rate do not appear to match | Internal error detected | Print ECG and calculate the pace rate. |
| Improper sensing (for example, sensing on T-waves) | QRS complex too small | Select another lead. |
| | T-wave too large | Adjust ECG size. |
| - | | |

Chapter 5 | Therapy

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---------------------------------|--|---|
| SYNC mode will not activate | PACER is on. Pacing and Sync are separate functions and are not allowed at the same time. | Discontinue pacing, if appropriate for the patient, and press SYNC. |
| Defibrillator will not turn off | Pacemaker is on | Turn off PACER and then press and hold ON for at least 2 seconds. |

For general troubleshooting tips, see General Troubleshooting Tips (on page 214).

Pediatric ECG Monitoring and Manual Mode Therapy Procedures

WARNING

Possible Patient Skin Burns

Do not use pediatric QUIK-COMBO electrodes on adults or larger children. Delivery of defibrillation energies equal to or greater than 100 joules (typically used on adults) through these smaller electrodes increases the possibility of skin burns.

WARNING

Possible Pediatric Patient Skin Burns

Noninvasive pacing may cause patient skin irritation and burns, especially with higher pacing current levels. Inspect underlying skin of the ♥ electrode frequently after 30 minutes of continuous pacing. Discontinue noninvasive pacing if skin burn develops and another method of pacing is available. On cessation of pacing, immediately remove or replace electrodes with new ones.

For pediatric patients, follow the procedures for ECG monitoring, manual defibrillation, synchronized cardioversion, and pacing except for the following:

- Use the appropriate paddle accessory based on the weight of the child.
- Select the appropriate defibrillation energy for the weight of the child according to the American Heart Association (AHA) recommendations or local protocol. Using energy levels of 100 joules or greater is likely to cause burns.
- When pacing, inspect the patient's skin under the heart electrode frequently for signs of burns.

Note: The amount of pacing current needed for capture is similar to the pacing current needed for adults. For more information about pediatric paddles and electrodes, see Paddle Accessory Options (on page 149).

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Chapter 6

Paddle Accessory Options

This chapter provides information about the paddle accessory options that may be used with the LIFEPAK 15 monitor/defibrillator.

| QUIK-COMBO Therapy Electrodes | 151 |
|--|-----|
| Standard Paddles | 154 |
| Sterilizable Internal Defibrillation Paddles | 158 |

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QUIK-COMBO Therapy Electrodes

Physio-Control QUIK-COMBO therapy electrodes are pre-gelled, self-adhesive therapy electrodes used for defibrillation, synchronized cardioversion, ECG monitoring, and pacing.



Figure 35 QUIK-COMBO Therapy Electrodes

A QUIK-COMBO therapy electrode set:

- Is a substitute for standard paddles.
- Provides Lead II monitoring signal when placed in the anterior-lateral position.
- Quickly restores the ECG trace on the monitor following defibrillation.

Always have immediate access to a spare set of therapy electrodes.

To help prevent therapy electrode damage:

- Only open electrode package immediately prior to use.
- Slowly peel back the protective liner on the electrodes, beginning with the cable connection end.
- Do not trim therapy electrodes.
- Do not crush, fold, or store the electrodes under heavy objects.
- Store therapy electrodes in a location where temperatures are between 15° and 35°C (59° and 95°F). Continuous exposure to the higher temperatures within this range will shorten the life of the electrodes.

Several types of QUIK-COMBO therapy electrodes are available as described in Table, QUIK-COMBO Electrodes (on page 152).

IMPORTANT! Infant/Child Reduced Energy Defibrillation Electrodes are not compatible with the LIFEPAK 15 monitor/defibrillator.

QUIK-COMBO Therapy Electrodes

Table 25 QUIK-COMBO Electrodes

| TYPE | DESCRIPTION |
|---|--|
| QUIK-COMBO | Electrodes, with 61 cm (2 ft) of lead wire, designed for patients weighing 15 kg (33 lb) or more |
| QUIK-COMBO RTS | Electrodes, providing a radio-transparent electrode and lead wire set, designed for patients weighing 15 kg (33 lb) or more |
| QUIK-COMBO with REDI-PAK® preconnect system | Electrodes designed for patients weighing 15 kg (33 lb) or more and that allow preconnection of the electrode set to the device while maintaining electrode shelf life and integrity |
| Pediatric QUIK-COMBO RTS | Electrodes designed for patients weighing 15 kg (33 lb) or less |

Connecting Therapy Electrodes

To connect QUIK-COMBO therapy electrodes to the QUIK-COMBO therapy cable:

- 1. Open the protective cover on the therapy cable connector (see the following figure).
- 2. To insert the QUIK-COMBO electrode connector into the therapy cable connector, align the arrows and press the connectors firmly together.



Figure 36 Connect QUIK-COMBO Electrodes to Therapy Cable

Replacing and Removing Therapy Electrodes

Replace adult QUIK-COMBO electrodes with new electrodes after one of the following occurs:

- 50 defibrillation shocks
- 24 hours on the patient's skin
- · 8 hours of continuous pacing

Replace pediatric QUIK-COMBO electrodes with new electrodes after one of the following occurs:

- 25 defibrillation shocks
- 24 hours on the patient's skin
- · 8 hours of continuous pacing

To remove QUIK-COMBO therapy electrodes from the patient:

1. Slowly peel back the therapy electrode from the edge, supporting the skin as shown.



Figure 37 Removing Therapy Electrodes from Skin

- 2. Clean and dry the patient's skin.
- 3. When applying new electrodes, adjust the positions slightly to help prevent skin burns.
- 4. Close the protective cover on the therapy cable connector when the cable is not in use.

Cleaning and Inspection

QUIK-COMBO electrodes are not sterile or sterilizable. They are disposable and are for a single patient application. Do not autoclave, gas sterilize, immerse in fluids, or clean electrodes with alcohol or solvents.

Include daily inspection of the QUIK-COMBO therapy electrode package as part of your defibrillator test routine. Daily inspection helps ensure that the therapy electrode has not exceeded the electrode package Use By date and is ready for use when needed. For more information about daily inspection and testing, see the Operator's Checklist in the back of this manual.

Standard Paddles

Adult Standard Paddles

Standard paddles are hard, hand-held paddles that are applied to the patient's chest to briefly monitor the ECG or to deliver defibrillation shocks. The following figure describes the features of the standard paddles.

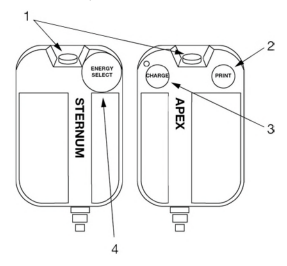


Figure 38 Standard Paddles

FIGURE LEGEND

- 1 (Shock) buttons. Discharge the defibrillator. Both buttons must be pressed simultaneously to deliver energy.
- 2 **PRINT** button. Activates printer. Function is identical to **PRINT** button on front panel.
- 3 **CHARGE** button. Charges the defibrillator. Adjacent **CHARGE** indicator flashes when device is charging and glows steadily when fully charged.
- 4 **ENERGY SELECT** dial. Rotary dial changes energy levels displayed on the screen.

A standard paddle set:

- Can be used instead of QUIK-COMBO therapy electrodes.
- Provides Lead II monitoring signal when held in the anterior-lateral position.
- Is used for defibrillation, synchronized cardioversion, and QUIK-LOOK® ECG checks.

To help prevent standard paddles damage:

- Handle with care to prevent damage to paddle surfaces.
- Store in paddle wells on the device to protect the electrode surface.
- Clean dried or wet gel from the electrode surface after each use.

Cleaning and Inspecting Standard Paddles

After each use:

- 1. Wipe standard paddle electrodes, handles, paddle wells, cables, and connector with mild disinfectant or soap and water solution. Do not immerse or soak.
- 2. Dry thoroughly.
- 3. Examine paddle surfaces, handles, cables, and connectors for damage or signs of wear.
 - Cables that show signs of wear such as loose cable connections, exposed wires, or cable connector corrosion must be removed from use immediately.
 - Paddles that have rough or pitted electrodes should be removed from use immediately.

Note: Standard paddles are not sterile or sterilizable. Do not autoclave, gas sterilize, immerse in fluids, or clean with alcohol or solvents.

Testing Standard Paddles

Include inspecting and testing of the standard paddles as part of your defibrillator test routine. Daily inspection and testing helps ensure that the standard paddles are in good operating condition and are ready for use when needed. For more information about inspection and testing, see the Operator's Checklist in the back of this manual.

Pediatric Paddles

Pediatric paddles slide onto adult paddles. Pediatric paddles should be used for patients weighing less than 10 kg (22 lb) or for patients whose chest size cannot accommodate the adult hard paddles.

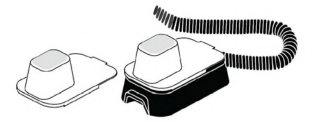


Figure 39 Pediatric Paddles

Use the adult paddle controls for selecting energy and charging. Each pediatric paddle attachment has a metal spring plate with a contact on it that transfers defibrillation energy from the adult paddle electrode to the pediatric paddle. This solid cadmium-silver contact will not scratch the adult paddle electrode.

Note: Inspect the spring plates and the contacts routinely to make sure that they are clean and intact.

Attaching Pediatric Paddles

To attach the pediatric paddles:

- 1. Slide the paddles onto clean adult paddles, starting at the front of the adult paddle (see following figure).
- 2. Slide the pediatric paddle until you feel the paddles lock in place.

Note: Do not use conductive gel between adult and pediatric paddles.

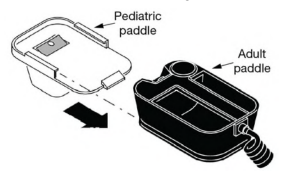


Figure 40 Attaching a Pediatric Paddle

Removing Pediatric Paddles

To remove pediatric paddles:

- 1. Press down on the rear tab.
- 2. Slide the pediatric paddle off.



Figure 41 Removing a Pediatric Paddle

Placing Pediatric Paddles

Adult paddles are recommended if the paddles fit completely on the child's chest. Allow at least 2.5 cm (1 in.) of space between the paddles.

For infants with very small chests, pediatric paddles may be too large to place in the anterior-lateral position. In this situation, place paddles in the anterior-posterior position. Holding the paddles against the chest and back supports the patient on his or her side.

Do not use the pediatric paddles on adults or older children. Delivery of recommended adult energies through this relatively small electrode surface increases the possibility of skin burns.

Anterior-Lateral Placement. Standard pediatric paddle placement includes (see following figure):

- **STERNUM** paddle to the patient's right upper torso, lateral to the sternum and below the clavicle.
- **APEX** paddle lateral to the patient's left nipple in the midaxillary line, with the center of the paddle in the midaxillary line, if possible.

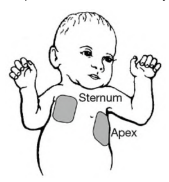


Figure 42 Anterior-Lateral Paddle Position

Anterior-Posterior Placement. Place the **STERNUM** paddle anteriorly over the left precordium and the **APEX** paddle posteriorly behind the heart in the infrascapular area (see following figure).

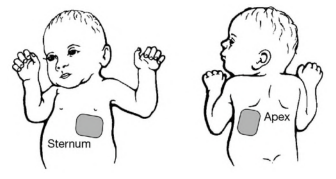


Figure 43 Anterior-Posterior Paddle Position

Cleaning and Inspecting Pediatric Paddles

Individually protect paddles before and after cleaning to prevent damage to paddle surfaces. After each use:

- Wipe or rinse paddle electrodes, cable connector, paddle handles, and cables with mild soap and water or disinfectant using a damp sponge, towel, or brush. Do not immerse or soak.
- 2. Dry thoroughly.
- 3. Examine paddle surfaces, connector, handles, and cables for damage or signs of wear.
 - Cables that show signs of wear such as loose cable connections, exposed wires, or cable connector corrosion should be removed from use immediately.
 - Paddles that have rough or pitted electrodes should be removed from use immediately.

Sterilizable Internal Defibrillation Paddles

Physio-Control internal paddles are specifically designed for open chest cardiac defibrillation.



Figure 44 Sterilizable Internal Defibrillation Paddles

Internal paddles are available in several sizes. To order internal paddles, contact your Physio-Control representative. In the USA, call Customer Support at 1.800.442.1142, option 2.

For complete information about using internal paddles to provide open chest cardiac defibrillation, see the *Instructions for Use* provided with the internal paddles.

Chapter 7

Data Management

This chapter describes how to manage current and archived Patient Records when using the LIFEPAK 15 monitor/defibrillator.

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Patient Records and Reports

When you turn on the LIFEPAK 15 monitor/defibrillator, a new Patient Record is created and stamped with the current date and time. All events and associated waveforms are digitally stored in the Patient Record as reports, which you can print, transmit, or download to the LIFENET® System, or to post-event review products such as CODE-STATTM or DT EXPRESSTM software. For information on how to print a report, see How to Print a Current Report (on page 167). For information on how to transmit or download a report, see Data Transmission (on page 173). When you turn off the device, the current Patient Record is saved in the archives.

You can also print, transmit, download, or delete any Patient Records that are stored in the archives. To access the archives, press **OPTIONS** and then select **ARCHIVES**. When you enter Archive mode, patient monitoring ends and the current Patient Record is saved and closed. Turn off the device to exit Archive mode. For more information, see Managing Archived Patient Records (on page 169).

Report Types

The reports that are available in a Patient Record depend on the features in your device and how your device is set up. For information on setting up your device, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device. The following table describes the various report types that may exist in a Patient Record and how they can be accessed.

Table 26 Report Types

| REPORT TYPE | DESCRIPTION | PRINT FROM MONITOR | TRANSMIT |
|--|---|--------------------|------------|
| 12-Lead ECG Report | The diagnostic 12-lead ECG report. For more information, see Printed 12-Lead ECG Report Formats (on page 63). | X | X ¹ |
| CODE SUMMARY™ Critical Event Record | Includes patient information, event and vital sign log, and waveforms associated with events (for example, defibrillation). For more information, see CODE SUMMARY Report (on page 162). | Х | X |
| Vital Signs Summary | Includes patient information and event and vital sign log. | Х | Х |
| Trend Summary | Includes patient information, vital sign log, and vital sign graphs. | X | Х |
| Snapshot Report | Includes patient information and 8 seconds of waveform data captured at the time of transmission. | | Х |
| Continuous Report ² | Provides real-time waveform data, acquired when the device is powered on and electrodes are connected or other waveform data is displayed in channels 2 or 3. Only for post-event review with CODE-STAT or DT EXPRESS software. | | Х |

¹ Transmission of a 12-lead ECG report automatically includes transmission of the Vital Signs Summary.

² To obtain CPR analytics using CODE-STAT software, the patient's ECG must be monitored using **PADDLES** lead in Channel 1.

Patient Records and Reports

Note: All reports that are transmitted to the LIFENET System include the following information:

- Battery status
- Power adapter status
- · Device usage information
- · Manufacturing configuration settings
- 3:00 A.M. self-test results

CODE SUMMARY Report

The LIFEPAK 15 monitor/defibrillator automatically stores a CODE SUMMARY report as part of the Patient Record for each patient. The CODE SUMMARY report can be set up to always print in a particular format. The available formats are shown in the following table. For CODE SUMMARY setup information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

To generate a CODE SUMMARY report, press **CODE SUMMARY**. If you interrupt printing of a CODE SUMMARY report, the entire CODE SUMMARY report is reprinted when printing is resumed. "Code Summary Complete" prints immediately following the last waveform event.

Table 27 CODE SUMMARY Formats

| FORMAT | ATTRIBUTES |
|---------------|--|
| Long format | Preamble |
| | Event/vital sign log |
| | Event waveforms |
| | 12-lead ECG reports |
| | Trend Summary |
| Medium format | Preamble |
| | Event/vital sign log |
| | Event waveforms |
| | Trend Summary |
| Short format | Preamble |
| | Event/vital sign log |
| | Trend Summary |

Note: When CODE SUMMARY reports are transmitted, they are always sent in the long format. Transmitted CODE SUMMARY reports do not include the Trend Summary.

The CODE SUMMARY report always contains the Preamble and the Event/Vital Sign Log. See the following figure for an example.

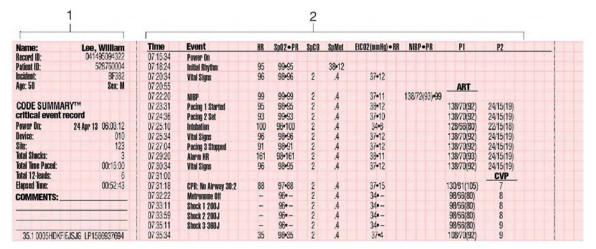


Figure 45 CODE SUMMARY Report

FIGURE LEGEND

- 1 Preamble
- 2 Event/Vital Sign Log

Preamble

The preamble consists of patient information (name, patient ID, age, and sex) and device information (date, time, and therapy information) as shown in the preceding figure. The defibrillator automatically enters a unique identifier in the ID field for each Patient Record. This identifier is composed of the date and time that the defibrillator is turned on. The Incident field allows you to enter up to 14 alpha-numeric characters to link the device to other documents such as an EMS Run Report.

Event/Vital Sign Log

The LIFEPAK 15 monitor/defibrillator documents events and vital signs in chronological order. Events are operator or device actions, such as actions that are related to monitoring, pacing, AED therapy, or data transmission. Values for each active vital sign are entered into the log automatically every 5 minutes and for each event. The following table lists events that may be found in the Event Log.

Patient Records and Reports

Table 28 Possible Event Log Entries

Monitoring

- Check patient
- Initial rhythm
- Replace battery
- 12-lead
- NIBP
- Alarm events
- IP label change
- Vital signs
- 5-wire on/off
- SpCO/SpMet Advisory

AED

- Connect electrodes
- Motion
- Analysis
- Analysis stopped
- Shock advised
- No shock advised

CPR Metronome

- On/Off
- Age-Airway changed

Defibrillation

- Manual mode
- · Charge removed
- Shock X, XXXJ
- Shock X, Abnormal

Operator Initiated

- Event
- Alarms on/off
- Print
- VF/VT alarm on/off
- Sync on/off
- Snapshot
- Internal pacer detection on/off

Pacing

- Started
- Set
- Changed
- Stopped
- Paused

Transmission

- Transmission complete
- Transmission failed
- Transmission cancelled

Memory Status

- Out of waveform memory (memory low)
- Out of event memory (memory full)

Waveform Events

In addition to being documented in the Event Log, therapy and other selected events also capture waveform data that are printed with the long and medium CODE SUMMARY report. The waveform events and the characteristics of waveform data are described in the following table.

Table 29 Waveform Events

| EVENT NAME | WAVEFORM DATA (WHEN CAPTURED) |
|---------------------------|---|
| INITIAL RHYTHM | 8 seconds after leads on |
| CHECK PATIENT | 8 seconds prior to alert |
| SHOCK or NO SHOCK ADVISED | 2-3 segments of analyzed ECG. Each segment is 2.7 seconds |
| ANALYSIS X STOPPED | 8 seconds of data prior to cessation of analysis |
| SHOCK X | 3 seconds prior to shock and 5 seconds after shock |
| PACING X STARTED | 8 seconds prior to increase of current from 0 |
| PACING X SET | 8 seconds after ppm and mA are stable for 10 seconds |
| PACING X CHANGED | 8 seconds after pacing rate, current, or mode is changed |
| PACING X STOPPED | 3 seconds prior to pacing current is zero and 5 seconds after |

Chapter 7 | Data Management

| EVENT NAME | WAVEFORM DATA (WHEN CAPTURED) |
|-----------------|---|
| PACING X PAUSED | Initial 8 seconds while PAUSE is pressed |
| ALARM* | 3 seconds prior to violated parameter and 5 seconds after |
| EVENT* | 3 seconds prior to event selection and 5 seconds after |
| PRINT | 3 seconds prior to pressing PRINT and 5 seconds after |
| 12-LEAD | 10 seconds after 12-LEAD is pressed |
| SNAPSHOT | 3 seconds prior to and 5 seconds after SNAPSHOT requested |
| VITAL SIGNS | 3 seconds prior to and 5 seconds after vital signs are acquired |

^{*}To reduce the length of the CODE SUMMARY report, storing waveform data with these events can be set to OFF (see the LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device).

Waveform events are preceded by a header that includes the following information:

Patient data

Vital signs

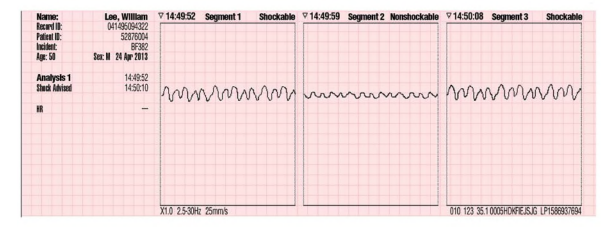
Event name

Device configuration information

Therapy data*

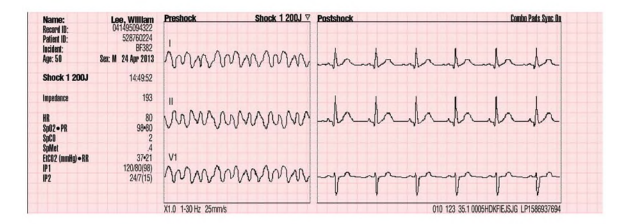
The following figures show four examples of waveform events as they would appear in the CODE SUMMARY report.

Analysis Event

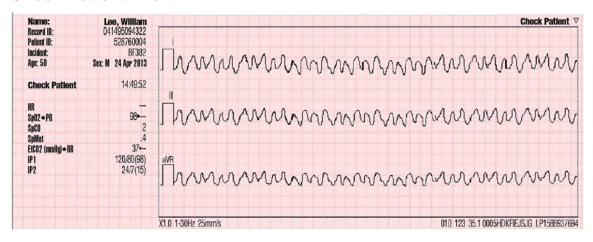


^{*}Patient impedance (in ohms) appears on shock reports when using disposable defibrillation electrodes. This impedance is measured just prior to the shock and is used to determine voltage compensation.

Shock Event



Check Patient Event



Pacing Event



Figure 46 Waveform Event Printout Examples

Memory Capacity

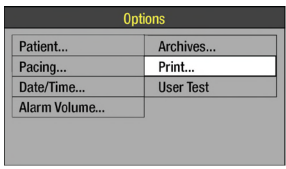
The LIFEPAK 15 monitor/defibrillator retains data for two or more patients when you switch power off or remove the batteries. The number of patient reports that the LIFEPAK 15 monitor/defibrillator can store depends on various factors, including the number of displayed waveforms, the duration of each use, and the type of therapy. The total capacity is 360 minutes of continuous ECG, 90 minutes of continuous data from all channels, or 400 single waveform events. The maximum memory capacity for a single patient includes up to 200 single waveform reports and 90 minutes of continuous ECG. When the defibrillator reaches the limits of its memory capacity, the defibrillator deletes an entire Patient Record using a "first in, first out" priority to accommodate a new Patient Record. Deleted Patient Records cannot be retrieved.

Managing Current Patient Records

You can add specific patient information to a current Patient Record. For more information, see Entering Patient Data (on page 42).

How to Print a Current Report

To print a current report:



- 1 Press **OPTIONS**. The Options menu appears.
- 2 Select **PRINT**. The Options/Print menu appears.

Managing Current Patient Records

| | otions / Print |
|--------|----------------|
| Print | |
| Report | Code Summary |
| Format | 3-Channel |
| Mode | Monitor |
| Speed | 25mm/sec |

3 If the REPORT, FORMAT, and MODE settings are correct, select PRINT. Otherwise, make changes as desired.

Select a REPORT:

- CODE SUMMARY
- TREND SUMMARY
- VITAL SIGNS
- 12-LEAD

Note: A check next to a 12-lead report indicates that the report was previously printed.

Select a **FORMAT** (for 12-Lead ECG only):

- 3-CHANNEL
- 4-CHANNEL

Select a **MODE** to change the frequency response of ECG reports:

- MONITOR
- DIAGNOSTIC (12-Lead reports always print in Diagnostic mode)

Select the **SPEED** option on this menu to change the speed of the continuous printout when the **PRINT** button is pressed. Note that this **SPEED** option does not affect reports that are printed from this menu. Available printing speeds for the **PRINT** button are:

- 12.5 mm/sec
- 25 mm/sec

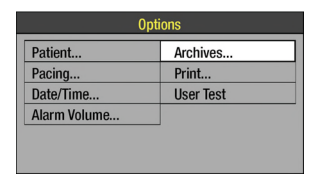
Managing Archived Patient Records

When you turn off the LIFEPAK 15 monitor/defibrillator, the current Patient Record is saved in the archives. You can print, edit, delete, or download archived records. For information about downloading to CODE-STAT software, see Data Transmission (on page 173). You can also transmit individual reports from an archived Patient Record. For information about transmitting an archived report, see Data Transmission (on page 173).

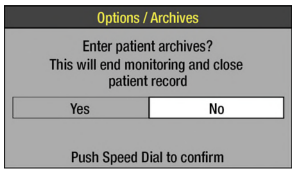
Note: When you enter Archive mode, patient monitoring ends (for example, no ECG, no alarms) and the current Patient Record is saved and closed.

Accessing Archive Mode

To enter Archive mode:



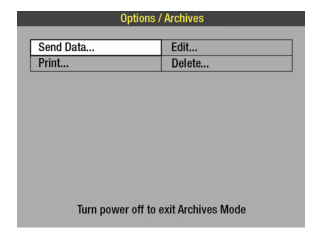
- Press **OPTIONS**. The Options menu appears.
- 2 Select **ARCHIVES**. The Options/Archives menu appears.



3 Select YES. The device enters Archive mode and the Options/Archives menu appears. Note: Your device may be set up so that you must enter a password to enter Archive mode.

Note: To exit Archive mode, power off the device.

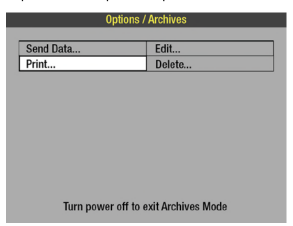
Managing Archived Patient Records



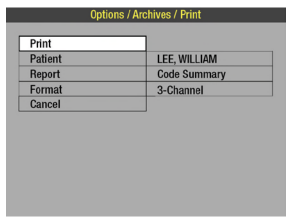
You can send, print, edit, or delete an archived record. For information about sending an archived record, see Data Transmission (on page 173).

Printing Archived Patient Reports

To print archived patient reports:



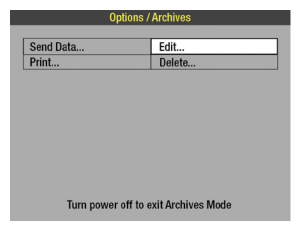
1 In Archive mode, select PRINT. The Options/ Archives/Print menu appears showing the current patient.



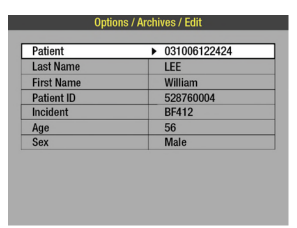
- 2 If the **PATIENT**, **REPORT**, and **FORMAT** settings are correct, go to Step 6.
- 3 To select a different patient, select PATIENT and then select the desired patient from the list.
- 4 To select a different report, select **REPORT** and then select one of the following:
 - CODE SUMMARY
 - TREND SUMMARY
 - VITAL SIGNS
 - 12-LEAD
- 5 To select a different format, select **FORMAT** and then select one of the following (for 12-Lead ECG only):
 - 3-CHANNEL
 - 4-CHANNEL
- 6 Select **PRINT**. The archived report is printed.

Editing Archived Patient Records

To edit archived patient records:



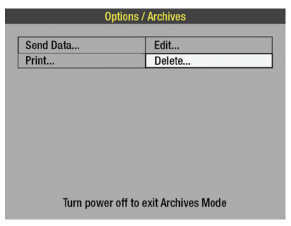
1 In Archive mode, select **EDIT**. The Options/Archives/Edit menu appears.



- 2 Select PATIENT.
- 3 Add the necessary patient information. Only blank fields may be edited.
- 4 Press **HOME SCREEN** and then turn off the device.

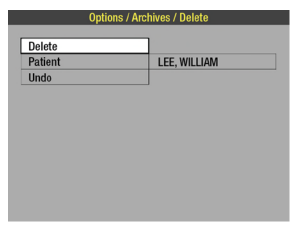
Deleting Archived Patient Records

To delete archived patient records:



1 In Archive mode, select **DELETE**. The Options/Archives/Delete menu appears.

Managing Archived Patient Records



- 2 To permanently remove the Patient Record that is displayed, select **DELETE**.
- 3 To see the list of all patient records, select **PATIENT**. The patient list appears. Select the Patient Record you want to delete.
- 4 To undo the delete operation, immediately select **UNDO**. If you continue with other device operations, you cannot undo the deletion.
- 5 Press **HOME SCREEN** and then turn off the device.

Chapter 8

Data Transmission

This chapter describes how to transmit Patient Records and reports from the LIFEPAK 15 monitor/defibrillator.

| About Transmitting Patient Records and Reports | 175 |
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| Preparing the Monitor for Transmission | 176 |
| Using Bluetooth Wireless Communication | 176 |
| Using a Direct Connection | 181 |
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About Transmitting Patient Records and Reports

You can transmit current and archived data from the LIFEPAK 15 monitor/defibrillator to the LIFENET® System or to post-event review products such as CODE-STAT™ or DT EXPRESS™ software.

The LIFEPAK 15 monitor can transmit patient reports using the following methods:

- Bluetooth® wireless connection—If your LIFEPAK 15 monitor has the Bluetooth feature installed and enabled, you can transmit data using a wireless connection.
- Direct cable connection—You can use a special cable to establish a direct connection from the LIFEPAK 15 monitor to a PC or gateway, and transmit data using this wired connection.

The following figure represents an overview of the data transmission process.

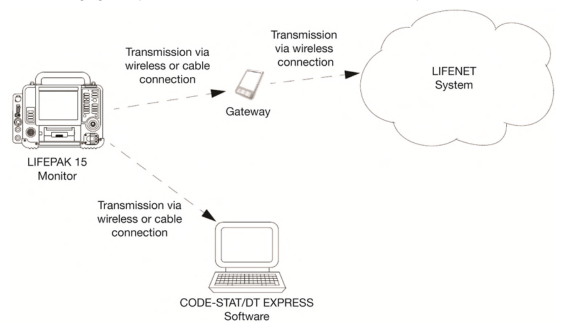


Figure 47 Transmitting Data from the LIFEPAK 15 Monitor/Defibrillator

For information about configuring your LIFEPAK 15 monitor to work in the LIFENET System, see the LIFENET System help documentation or contact your Physio-Control representative.

Preparing the Monitor for Transmission

Before you can transmit using a wireless or direct connection, you must define transmission sites and output ports in the LIFEPAK 15 monitor Setup mode.

For each transmission site, select an output port:

- For wireless transmission, set OUTPUT PORT to BLUETOOTH WIRELESS.
- For a direct connection, set **OUTPUT PORT** to **DIRECT CONNECT**.
- Set OUTPUT PORT to BOTH if you normally transmit using a Bluetooth connection but you
 need a direct cable backup. (If you set OUTPUT PORT to BOTH, make sure the Bluetooth LED
 is not illuminated before you attempt to transmit using a direct connection. The device will
 not transmit using the direct connection when a wireless connection is available.)

For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

Using Bluetooth Wireless Communication

Bluetooth technology is a short-range wireless communication technology that is available as an option on the LIFEPAK 15 monitor/defibrillator. When Bluetooth technology is installed, the Bluetooth icon appears on the Home Screen. See Figure, Bluetooth Icon on the Home Screen (on page 177).

See the *Bluetooth* label in battery well 2 for FCC and Industry Canada radio identification numbers.

A *Bluetooth* connection between the LIFEPAK 15 monitor and a target device is always initiated from the LIFEPAK 15 monitor.

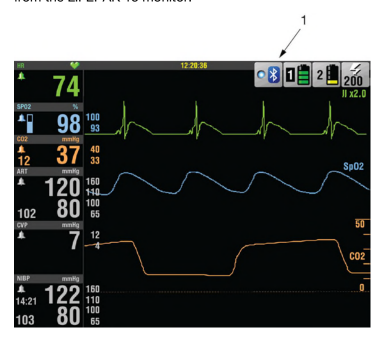


Figure 48 Bluetooth Icon on the Home Screen

FIGURE LEGEND

1 Bluetooth icon

The *Bluetooth* icon shows the status of the wireless connectivity in the device.

Bluetooth Passcodes

The LIFEPAK 15 monitor has a *Bluetooth* passcode that you define.

To transmit from the LIFEPAK 15 monitor to a headless gateway (a device that has no user interface), the *Bluetooth* passcode that you enter in the LIFEPAK 15 monitor must match the *Bluetooth* passcode that is preconfigured in the gateway. For information about the *Bluetooth* passcode in the headless gateway, see the documentation that ships with the gateway, or consult your system administrator or equipment technician.

To transmit from the LIFEPAK 15 monitor to a PC, you need to set a *Bluetooth* passcode in the LIFEPAK 15 monitor, and then enter that passcode on the PC, if prompted.

Bluetooth Search Filter

A *Bluetooth*-enabled LIFEPAK 15 monitor may discover numerous *Bluetooth* devices that are within range. To help filter out extraneous devices and find the specific target device that you want to transmit to, Physio-Control developed the Physio Service Class (PSC).

The PSC is a prefix that you can add to the friendly name of your target devices. Then when you set the **SEARCH FILTER** to **ON** in the LIFEPAK 15 monitor, only target devices that have the PSC

Using Bluetooth Wireless Communication

prefix in their names appear in the list of discovered devices (if they are powered on and discoverable).

The various PSC prefixes correspond to LIFEPAK 15 monitor modes of operation. The following table lists the LIFEPAK 15 monitor modes and the service class and friendly name prefix that is discoverable in each mode. For example, when the LIFEPAK 15 monitor is in Archive mode and the filter is on, it can discover devices whose friendly names begin with A_ or B_.

Table 30 Physio Service Class Prefixes

| LIFEPAK 15 MONITOR/DEFIBRILLATOR MODE | SERVICE CLASS | FRIENDLY NAME PREFIX |
|---|-------------------------------|-------------------------|
| LIFEPAK 15 monitor must be in Archive mode | Archive | A_ |
| LIFEPAK 15 monitor can be in AED, Manual, or Archive mode | Both Cardiac Care and Archive | В_ |
| LIFEPAK 15 monitor can be in AED or Manual mode | Cardiac Care | C_ |

For information about configuring the friendly name in your target devices, see the documentation provided with those devices.

Bluetooth Setup

Use the Bluetooth Setup menu to set up the Bluetooth transmission on the LIFEPAK 15 monitor.

To access the Bluetooth Setup menu:

| Blu | etooth Setup |
|---------------|---------------------|
| Connect | ► (Not Connected) |
| Search Filter | 0n |
| Passcode | 0000 |
| Wireless | 0n |
| Disconnect | |
| LIFEPAK 15 | Device ID: LP151234 |

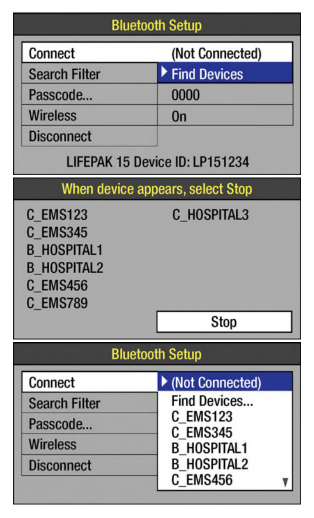
- On the HOME SCREEN, rotate the SPEED DIAL to outline the Bluetooth icon.
- 2. Press the **SPEED DIAL**. The *Bluetooth* Setup menu appears.
- Set SEARCH FILTER to ON if you want to find only devices that include the PSC in their friendly name; otherwise, set SEARCH FILTER to OFF.
- 4. Set a Bluetooth passcode.
 - To transmit to a headless gateway, enter the passcode that is preconfigured in the gateway.
 - To transmit to a PC, you may need to enter a passcode or acknowledge the connection.
- 5. Ensure that WIRELESS is set to ON.

Note: The default setting for **WIRELESS** is **ON**, and the default setting for **SEARCH FILTER** is **OFF**. Use the **WIRELESS** setting to turn off the wireless signal when operating the LIFEPAK 15 monitor in an environment where transmission is not desirable.

Establishing a Bluetooth Connection

You must know the friendly name of the target device that you want to connect to.

To establish a *Bluetooth* connection:



- 1. On the LIFEPAK 15 monitor, use the **SPEED DIAL** to select the *Bluetooth* icon and access the *Bluetooth* Setup menu.
- Select CONNECT and then select FIND DEVICES. This will disconnect any existing connections.

Note: If the LIFEPAK 15 monitor is set to **WIRELESS OFF**, wireless status changes to **WIRELESS ON**.

- The Find Devices menu appears. The monitor begins searching for Bluetooth devices that are in the area and that meet the search filter criteria.
- Devices are displayed in the order found—the most recently found device appears at the top of the list.
- When the desired device appears, press the SPEED DIAL to select STOP and end the search. You return to the Bluetooth Setup menu.
- 4. Use the **SPEED DIAL** to scroll through the list and select the desired device.
- If you are connecting to a PC, you may be prompted to acknowledge the connection. Enter the passcode, if requested, and then accept the connection.
- When the connection is made, an alert tone sounds, the Bluetooth LED on the Home Screen is illuminated, and CONNECTED TO (DEVICE NAME) briefly appears in the message area.

After you establish a *Bluetooth* connection, you are ready to transmit patient data. Proceed to Transmitting Reports (on page 183).

Re-establishing a Bluetooth Connection

The LIFEPAK 15 monitor retains in its memory two last-connected devices, limited to one in each mode—one for cardiac care (AED or Manual mode) and one for Archive mode. When the LIFEPAK 15 monitor is powered on and the wireless feature is set to **WIRELESS ON**, the monitor automatically searches for the last connected device. If the last connected device in that mode is turned on and within range, a connection is established automatically. When a connection is established, the *Bluetooth* LED is illuminated and **CONNECTED TO (DEVICE NAME)** appears in the message area.

Note: If **RESET DEFAULTS** is selected in Setup mode, the *Bluetooth* passcode is not reset. However, connections to the last-connected devices are reset (terminated). To re-establish a connection, use **FIND DEVICES**.

Table 31 Bluetooth Status

| Table of Blacketti Ctar | |
|-------------------------|--|
| BLUETOOTH ICON | DESCRIPTION |
| • 🐉 | The <i>Bluetooth</i> LED is illuminated when the <i>Bluetooth</i> feature is enabled in this device and this device is connected to another <i>Bluetooth</i> -enabled device. |
| • 🖁 | The <i>Bluetooth</i> icon appears but the LED is not illuminated when the <i>Bluetooth</i> feature is enabled in this device, but this device is currently not connected to another <i>Bluetooth</i> -enabled device. |
| × | A red X appears when the <i>Bluetooth</i> feature is installed in this device, but wireless communication is currently set to OFF or there is a <i>Bluetooth</i> malfunction. See Troubleshooting Tips for Data Transmission (on page 186). |

Preparing for a Wireless Transmission

Before you can send wireless transmissions from the LIFEPAK 15 monitor, you must prepare the monitor and target devices for communication.

The target device must:

- Be Bluetooth-enabled, turned on, and discoverable.
- Have the LIFENET PC Gateway application or the patient care reporting software CODE-STAT or DT EXPRESS installed and running.
- Have a Bluetooth COM port configured for incoming data.
- Have an established friendly name.

The LIFEPAK 15 monitor must:

- Have at least one transmission site defined that has OUTPUT PORT set to BLUETOOTH WIRELESS.
- Have a *Bluetooth* passcode that matches the passcode in the target device, if the target device requires a passcode.
- Have SEARCH FILTER set to ON if you are using the Physio Service Class. For information about the Physio Service Class, see Bluetooth Search Filter (on page 177) later in this chapter.

Terminating a Bluetooth Connection

When the *Bluetooth* LED is illuminated, the LIFEPAK 15 monitor has a wireless connection established with another *Bluetooth* device.

To terminate a *Bluetooth* connection:

- 1. Use the SPEED DIAL to select the Bluetooth icon and access the Bluetooth Setup menu.
- 2. Select **DISCONNECT.** The *Bluetooth* connection is terminated and is not retained as the last connected device.

Using a Direct Connection

A special cable can be used to create a direct connection between the LIFEPAK 15 monitor and a gateway or PC. The following figure shows the equipment connections to send reports directly to a computer using a direct cable connection.

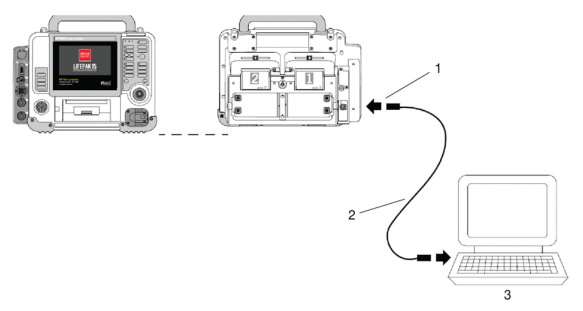


Figure 49 Data Transmission using a Direct Connection

FIGURE LEGEND

- System connector
- 2 LIFEPAK monitor to PC cable
- 3 Computer

WARNING

Shock Hazard

All equipment connected to the system connector must be battery powered or electrically isolated from AC power according to IEC 60601-1. If in doubt, disconnect the patient from the defibrillator before using the system connector. Only use Physio-Control recommended data transmission cables. For more information, contact Physio-Control Technical Support.

WARNING

Improper Device Performance Hazard

RF communication equipment such as cell phones, modems and radios may interfere with the performance of the monitor/defibrillator. If the monitor/defibrillator is used near RF communication equipment, observe the recommended separation distances in Recommended Separation Distances Between Portable and Mobile RF Communications Equipment and the LIFEPAK 15 Monitor/Defibrillator (on page 266). Certain RF communication equipment can be used at distances that are less than those recommended in these operating instructions. If the separation distance is less than the recommended distance, use only equipment recommended by Physio-Control and observe the monitor/defibrillator to verify normal operation.

To establish a direct connection:

1. Position the PC or LIFENET Gateway within reach of the LIFEPAK 15 monitor.

Note: If you are storing a LIFENET Gateway (modem) in the carrying case, only store the modem in the side pouch. Do not store LIFENET Gateways in the back pouch.

- Configure a COM port on the PC for incoming data.
- 3. Connect the cable to the system connector on the monitor and to the PC.
- 4. If using CODE-STAT or DT EXPRESS software, open the download wizard on the PC and select the LIFEPAK 15 monitor.

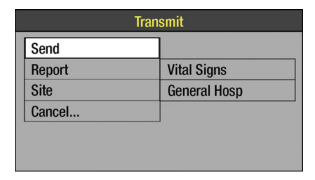
After you establish a direct connection, you are ready to transmit patient data. Proceed to Transmitting Reports (on page 183).

Transmitting Reports

After you have established a wireless or direct connection, you can transmit Patient Records and reports. All patient reports can be transmitted real time during patient monitoring (Manual or AED mode), or reports can be transmitted post event (Archive mode).

How to Transmit a Current Patient Report

To transmit a current patient report:

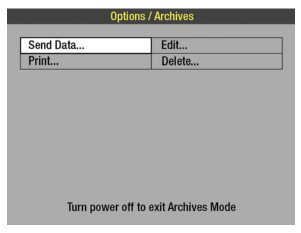


- 1. Press **TRANSMIT**. The Transmit menu appears.
- 2. Use the **SPEED DIAL** to select the desired **REPORT** and **SITE**, if necessary.
- Select SEND. The patient report is transmitted. The status of the transmission appears in the message area.

How to Transmit an Archived Patient Report

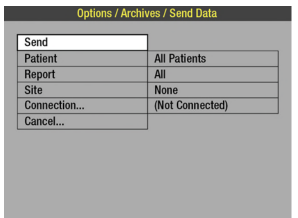
When you turn off the LIFEPAK 15 monitor/defibrillator, the current Patient Record is saved in the archives. For information about accessing Archive mode, see Data Management (on page 159).

To transmit an archived patient report:

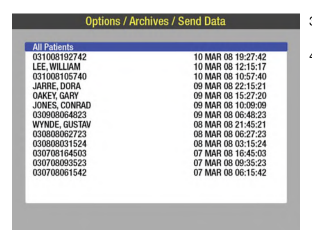


1. In the Options/Archives menu, select **SEND DATA**. The Options/Archives/Send Data menu appears.

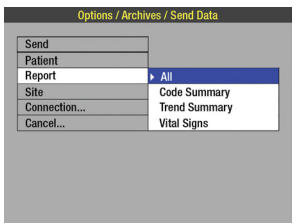
Transmitting Reports



2. If the **PATIENT**, **REPORT**, and **SITE** are correct, proceed to Step 7.



- To transmit records for a particular patient, select PATIENT. A list of patients appears.
- Select the patient.



- To transmit a specific report, select REPORT and then select the report.
- To select a transmission site, select SITE and then select the site. Make sure you specify a site whose OUTPUT PORT is configured for the transmission method you are using.
- To transmit using a wireless transaction, select CONNECTION and proceed with establishing a Bluetooth connection. For more information, see Establishing a Bluetooth Connection (on page 179).
- 8. Select **SEND**. The patient report is transmitted. The status of the transmission appears in the message area.

Transmission Status Report

Whenever you attempt to transmit a record, a transmission report is automatically printed at the completion of the transmission attempt. The transmission report indicates the date and time of the transmission attempt and the final status of the transmission.

Cancelling a Transmission

You can cancel a transmission that is in process. To cancel a transmission, select **CANCEL** on the Transmit menu if you are transmitting a current record, or select **CANCEL** on the Options/Archives/ Send Data menu if you are transmitting an archived record.

Considerations When Transmitting Data

When considering any treatment protocol that involves transmitting patient data, be aware of possible limitations. Successful transmission depends on access to public or private network services that may or may not always be available. This fact is especially true for wireless communication, which is influenced by many factors, such as:

- Geography
- Location
- Weather
- Number of wireless devices in the area

Treatment protocol must always take into account the fact that data transfer *cannot be assured* using wireless communication. Your treatment protocol must include contingency planning for interrupted data transmission.

Periodically test your device transmission function to ensure that the device and transmission accessories are ready for use.

Troubleshooting Tips

| Table 32 Troubleshooting | | |
|---|---|--|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
| Bluetooth icon on LIFEPAK 15 monitor has red X across it | WIRELESS is set to OFF in the Bluetooth Setup menu | Set WIRELESS to ON. If red X remains, Bluetooth module in LIFEPAK 15 monitor may be faulty. Contact qualified service representative. |
| | WIRELESS is set to OFF in the setup options, so the WIRELESS default is OFF each time the LIFEPAK 15 monitor is turned on | Change WIRELESS setup option. See LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device. |
| | | If red X remains, Bluetooth module in LIFEPAK 15 monitor may be faulty. Contact qualified service representative. |
| | Bluetooth module in LIFEPAK 15 monitor may be faulty | Contact qualified service representative. |
| Bluetooth LED is not illuminated | Target device is off or cannot communicate with the LIFEPAK 15 monitor | Confirm that target device is on and discoverable.See the operating instructions for |
| | | your target device. |
| | Bluetooth module in LIFEPAK 15 monitor may be faulty | If other troubleshooting is unsuccessful, contact qualified service representative. |
| LIFEPAK 15 monitor does not automatically connect to last connected device | Target device is off or cannot communicate with the LIFEPAK 15 monitor | Confirm that target device is on and discoverable. |
| | Last connection to target device may have occurred when the LIFEPAK 15 monitor was in a different mode | Confirm that OUTPUT PORT is set to BLUETOOTH WIRELESS. Select FIND DEVICES and establish a new connection. |
| Device does not connect to last connected device after WIRELESS is set to ON | Bluetooth menu is displayed, which prevents discovery of devices | Press HOME SCREEN to exit menu and allow LIFEPAK 15 monitor to find last connected device. |
| UNABLE TO CONNECT message appears | LIFEPAK 15 monitor cannot establish wireless connection. Target device may not have the necessary software application or cannot accept data. | Verify target device is ready to receive transmissions. Attempt to retransmit. |
| Unable to find a particular <i>Bluetooth</i> device, or BLUETOOTH | Search filter may be on and target device does not have a PSC prefix | Confirm that target device is on and discoverable.Confirm friendly name of target |
| DEVICE NOT FOUND message appears | | device. Set SEARCH FILTER to OFF and then select FIND DEVICES again. |
| | | |

| ODCEDVATION . | DOCCIDI E CALLOE | CORRECTIVE A CTION |
|---|--|---|
| OBSERVATION | POSSIBLE CAUSE Target device is not functioning | CORRECTIVE ACTION |
| | Target device is not functioning | Confirm that target device is on and discoverable. |
| | | Confirm friendly name of target device. |
| | | If message still appears, contact the service provider for your target device. |
| | Bluetooth module in LIFEPAK 15 monitor may be faulty | Contact qualified service representative. |
| Unable to transmit data for post-event review using either direct connection or <i>Bluetooth</i> connection | Post-event review software is not installed on target device | Install CODE-STAT or DT EXPRESS post-event review software on target device. |
| | Post-event review software is not open and running on target device | Make sure the target device is running Device Communications or the download wizard. |
| | COM port is not configured for incoming data on target device | Configure COM port on target device. |
| | LIFEPAK 15 monitor not selected in download wizard on target device | Open download wizard on target device and select the LIFEPAK 15 monitor. |
| BLUETOOTH UNAVAILABLE message | Bluetooth module in LIFEPAK 15 monitor not | Turn LIFEPAK 15 monitor off and back on. |
| appears | responding | If message still appears, Bluetooth module may be faulty. Contact qualified service representative. |
| BLUETOOTH DEVICE NOT FOUND message | Unable to locate Bluetooth device | Verify target device is ready to receive transmissions. |
| appears | | Set SEARCH FILTER to OFF and then select FIND DEVICES again. |
| UNKNOWN DEVICE message appears | Bluetooth name discovery failed or timed out before the device name was obtained | Verify name of target device.Verify target device is ready to receive transmissions. |
| | | Attempt to retransmit. |
| Unable to transmit using a gateway device that has a functioning direct connection or <i>Bluetooth</i> connection | Transmission sites are not set up in LIFEPAK 15 monitor | Define transmission sites. Each site name must exactly match the name of the target device. See LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device. |
| | Transmission site names in LIFENET System do not match site names in LIFEPAK 15 monitor | Check site names in LIFENET System. |
| | Cellular communication is not working between the gateway and transmission sites | Use alternate method to communicate patient data. |
| | | |

Troubleshooting Tips

| OBSERVATION UNABLE TO TRANSMIT message appears | POSSIBLE CAUSE The LIFEPAK 15 monitor cannot connect to the device name selected The output port on the | Verify target device is ready to receive transmissions. Verify target device setup. Attempt to retransmit. Make sure the transmission site |
|--|---|---|
| | LIFEPAK 15 monitor is not configured for the transmission method you are using | OUTPUT PORT is configured for the type of transmission you are attempting. Attempt to retransmit. |
| | Target device unable to connect or unable to connect within timeout interval | Verify target device is ready to receive transmissions. Verify target device setup. Attempt to retransmit. |
| | The target device requires you to "accept" incoming communications | Check your target device for a required acknowledgment to connect. Enter passcode, when prompted. Set to "Always allow" if possible. |
| | Direct connection was disrupted | Attempt to retransmit.Verify cable connections.Attempt to retransmit. |
| TRANSMISSION FAILED message appears | Computer application program is not ready or is not available to receive transmission | Verify target device is running necessary software.Attempt to retransmit. |
| LOST DIRECT CONNECTION message appears | Direct connection was interrupted | Verify cable connections between LIFEPAK 15 monitor and gateway or PC. Attempt to retransmit. |
| LOST BLUETOOTH CONNECTION message appears | Connection with <i>Bluetooth</i> target device was interrupted | Verify target device is ready to receive transmissions.Attempt to retransmit. |
| TRANSMISSION CANCELLED message appears | Operator of the LIFEPAK 15 monitor cancelled transmission | Attempt to retransmit if cancelled in error. |

Chapter 9

Power Adapter

This section describes the AC Power Adapter and the DC Power Adapter.

| Basic Orientation | 191 |
|-------------------------|-----|
| Using the Power Adapter | 193 |
| General Maintenance | 196 |
| Warranty | 197 |

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Basic Orientation

The AC Power Adapter and DC Power Adapter are optional accessories for use only with the LIFEPAK 15 monitor/defibrillator. These power adapters:

- Provide operating power to the monitor/defibrillator with or without batteries installed.
- Provide power to charge batteries installed in the monitor/defibrillator.

The AC Power Adapter operates with either 120 or 240 Vac line power. The DC Power Adapter operates with 12 Vdc power. Installed batteries are charged whenever the power adapter is connected to the LIFEPAK 15 monitor/defibrillator. To help manage and maintain battery charge, the power adapter should be kept plugged into the defibrillator whenever possible. For more information about maintaining the batteries, see Battery Maintenance (on page 209).

Note: Although the monitor/defibrillator can operate using auxiliary power with no batteries installed, at least one battery should be installed at all times.

Note: If the monitor/defibrillator loses power for more than 30 seconds, it will revert to the user-configured default settings and begin a new patient record.

An optional output extension cable is available. The output extension cable is equipped with a breakaway connector to allow quick movement if needed. For more information about the breakaway feature, see Output Extension Cable with Breakaway Connector (on page 195).

IMPORTANT! Daily inspection and testing will help ensure that the power adapter is in good operating condition and is ready for use when needed. Refer to the LIFEPAK 15 monitor/defibrillator Operator's Checklist in the back of this manual.

Carefully read the *Power Adapter Instructions for Use* that are provided with the power adapter for complete instructions, warnings, cautions, and specifications.

WARNING

Possible Loss of Power During Patient Care

Physio-Control has no information regarding the performance or effectiveness of its LIFEPAK monitor/defibrillators if other manufacturers' power adapters are used. Using other manufacturers' power adapters may cause the device to perform improperly and invalidate the safety agency certifications. Use only power adapters that are labeled with the

LIFEPAK 15 device symbol shown here. 15



WARNING

Possible Loss of Power During Patient Care

Do not use the LIFEPAK 12 power adapter with the LIFEPAK 15 monitor/defibrillator. Use only power adapters that are labeled with the LIFEPAK 15 device symbol. 15

WARNING

Possible Loss of Power During Patient Care

If the monitor/defibrillator will be used in emergency environments that require battery power, the installed batteries must be kept fully charged. Keep the power adapter plugged into an auxiliary power source whenever possible to maintain the charge level.

WARNING

Possible Loss of Power During Patient Care

Do not connect more than one output extension cable between the power adapter and the defibrillator. The resultant voltage drop may prevent the power adapter from charging the batteries or operating the defibrillator. Always connect the power adapter directly to the defibrillator or use only one extension cable.

WARNING

Shock Hazard

Using a power line cord other than the one supplied with the power adapter could cause excess leakage currents. Use only the power line cord that is specified for use with the power adapter.

WARNING

Potential Performance Degradation

Physio-Control has no information regarding the performance or effectiveness of the LIFEPAK 15 monitor/defibrillator when the power adapter is used with a power inverter. It is the user's responsibility to verify that the monitor/defibrillator performs correctly when used with a power inverter.

WARNING

Possible Skin Injury

The power adapter may become warm when used for an extended period of time. Prolonged contact between exposed skin and a warm power adapter may cause skin irritation or burns. If a warm power adapter is placed against a patient, the operator should ensure that the patient's skin is adequately protected.



Figure 50 AC Power Adapter

Using the Power Adapter

This section provides information about operating the AC and DC power adapters that can be used with the LIFEPAK 15 monitor/defibrillator.

AC Power Adapter Operation

To use the AC Power Adapter:

- 1. Connect the AC power cord to the power adapter and a grounded AC outlet.
- 2. Verify that the green LED strip illuminates.
- 3. Connect the power adapter output cable to the power adapter.
- 4. Connect the green end of the power adapter output cable to the auxiliary power connector on the back of the monitor/defibrillator.
- 5. Verify that the **AUXILIARY POWER** LED on the defibrillator is illuminated.
- If at least one battery is installed in the device, verify that the BATTERY CHARGING indicator is illuminated or flashing. Indicator behaviors are shown in Table, Battery Charging Indicator Behaviors, below.

Table 33 Battery Charging Indicator Behaviors

| INDICATOR | DESCRIPTION |
|----------------|---|
| Steady green | Installed batteries are fully charged. |
| Flashing green | One or both installed batteries are being charged. |
| Off | No batteries are installed, or a battery is unable to be charged. |

7. Press the monitor/defibrillator **ON** button.

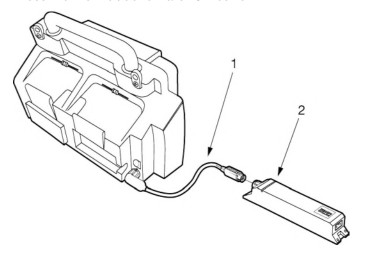


Figure 51 AC Power Adapter with LIFEPAK 15 Monitor/Defibrillator

FIGURE LEGEND

- 1 Power adapter output cable
- 2 LED strip

DC Power Adapter Operation

To use the DC Power Adapter:

- 1. Connect the DC power cable to the power adapter and a 12 Vdc power source.
- 2. Verify that the green LED strip illuminates.
- 3. Connect the power adapter output cable to the power adapter.
- 4. Connect the green end of the power adapter output cable to the auxiliary power connector on the back of the monitor/defibrillator.
- 5. Verify that the **AUXILIARY POWER** LED on the defibrillator is illuminated.
- 6. If at least one battery is installed in the device, verify that the **BATTERY CHARGING** indicator is illuminated or flashing. Indicator behaviors are shown in Table, Battery Charging Indicator Behaviors, below.

 Table 34 Battery Charging Indicator Behaviors

| INDICATOR | DESCRIPTION |
|----------------|--|
| Steady green | Installed batteries are fully charged. |
| | |
| Flashing green | One or both installed batteries are being charged. |

7. Press the defibrillator **ON** button.

Output Extension Cable with Breakaway Connector

One optional output extension cable may be connected between the power adapter and the power adapter output cable, if desired. The output extension cable is equipped with a breakaway connector that can be pulled apart without manually rotating the lock ring. With the breakaway connector, you can quickly separate the monitor/defibrillator from the power adapter without damaging the cables or connectors.

To use the breakaway feature, the power adapter and output extension cable must be secured as described in the *Power Adapter Instructions for Use*. The breakaway connector is designed to withstand routine breakaway use. However, to prolong the life of the connector, manually disconnect it whenever possible.

To order the output extension cable, contact your Physio-Control representative. In the USA, call Customer Support at 1.800.442.1142, option 2.

IMPORTANT! Do not use more than one output extension cable.

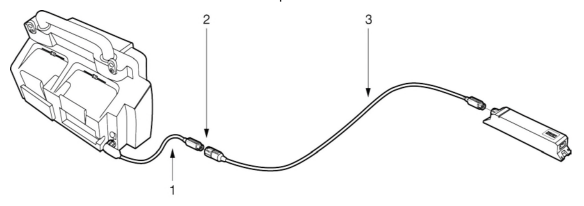


Figure 52 Output Extension Cable with Breakaway Connector

FIGURE LEGEND

- 1 Power adapter output cable
- 2 Breakaway connector
- 3 Output extension cable

General Maintenance

Maintenance and Service

The power adapter contains no serviceable parts. If the power adapter does not function correctly, contact your local Physio-Control representative for assistance.

Cleaning

WARNING

Possible Electrical Shock

Unplug the power adapter from the power source before cleaning.

CAUTION

Possible Equipment Damage

Do not clean any part of the power adapter or its accessories with phenolic compounds. Do not use abrasive or flammable cleaning agents. Do not attempt to sterilize this device or any accessories unless otherwise specified in accessory operating instructions.

To clean the power adapter:

- 1. Unplug the power adapter, if it is connected to an auxiliary power source.
- 2. Clean the power adapter, power cord, and cables with a damp sponge or cloth. Use only the cleaning agents listed below:
 - Quaternary ammonium compounds
 - Isopropyl alcohol
 - · Peracetic (peroxide) acid solutions
 - Sodium dichloroisocyanurate (NaDCC)
 - Chlorine bleach (1:10 dilution)

Note: Carefully clean the connector ports. Do not allow cleaning fluids to penetrate the exterior surfaces of the device.

Troubleshooting Tips

Table 35 Troubleshooting Tips for Power Adapter

| Table 35 Troubleshooting Tips | | | |
|---|---|---|--|
| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION | |
| POWER LED on power adapter does not light | Power cord not plugged into power adapter or power source | Connect power cord. | |
| | Defective power adapter or power cord | Replace with working power adapter and power cord. | |
| | Blown fuse or tripped circuit breaker in building | Contact qualified service personnel. | |
| AUXILIARY POWER LED on monitor/defibrillator not illuminated | Power adapter not properly connected to auxiliary power source or monitor/defibrillator | Check that power adapter is connected properly. | |
| | Defective power adapter or cables | Replace with working power adapter and cables. | |
| BATTERY CHARGING LED on monitor/defibrillator not illuminated | Power adapter not properly connected to auxiliary power source or monitor/defibrillator | Check that power adapter is connected properly. | |
| | Battery not properly inserted in battery well | Check that battery is properly inserted in battery well. | |
| | Unable to charge battery with power adapter because battery charge level is too low | Charge battery in Station-Mobile or REDI-CHARGE battery charger if available. Replace battery. | |
| | No batteries installed | Install at least one battery. | |
| | Defective battery | Remove battery from service and replace with working battery. | |
| | Unrecognized battery | Only use battery that is approved for use with the LIFEPAK 15 monitor/defibrillator. | |
| | Incompatible power adapter connected to the monitor/defibrillator | Only use power adapter that is approved for use with the LIFEPAK 15 monitor/defibrillator. | |
| | Defective power adapter or cables | Replace with working power adapter and cables. | |
| | Monitor/defibrillator unable to recognize installed battery | Contact qualified service personnel. | |

Warranty

To obtain a detailed warranty statement, contact your local Physio-Control representative or go to www.physio-control.com.

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Chapter 10

Maintaining the Equipment

This chapter describes how to perform operator-level maintenance, testing, and troubleshooting for the LIFEPAK 15 monitor/defibrillator and selected accessories. For additional information about accessories, refer to specific accessory operating instructions.

| General Maintenance and Testing | . 201 |
|---------------------------------|-------|
| Battery Maintenance | . 209 |
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General Maintenance and Testing

Periodic maintenance and testing of the LIFEPAK 15 monitor/defibrillator and accessories are important to help prevent and detect possible electrical and mechanical discrepancies. If testing reveals a possible discrepancy with the defibrillator or accessories, see General Troubleshooting Tips (on page 214). If the discrepancy cannot be corrected, immediately remove the LIFEPAK 15 monitor/defibrillator from service and contact a qualified service technician. For testing information regarding accessories, see the accessory operating instructions.

A **MAINTENANCE DUE** message can be set up to appear at selected intervals (3, 6, or 12 months) to remind you that the LIFEPAK 15 monitor/defibrillator is due for maintenance. The factory default is **OFF**, but it can be activated by service personnel.

An Operator's Checklist is included in the back of this manual. You may reproduce the checklist and use it to inspect and test the LIFEPAK 15 monitor/defibrillator. Daily inspection and test is recommended.

Maintenance and Testing Schedule

Table, Recommended Maintenance Schedule for Clinical Personnel (on page 202), lists the recommended maintenance and testing schedule. This schedule may be used in conjunction with the internal quality assurance program of the hospital, clinic, or emergency medical service where the defibrillator is used.

To ensure proper performance of the monitor/defibrillator, inspect and test the power adapter daily as described in the Operator's Checklist.

Cables and paddles are a critical part of therapy delivery and suffer wear and tear. Therapy cable testing as described in the Operator's Checklist is recommended on a daily basis. The Test Load ships with the device and is necessary for testing the QUIK-COMBO cable. Physio-Control recommends replacement of therapy cables every three years to reduce the possibility of failure during patient use.

The 12-lead ECG cable is a critical part of diagnosis and suffers wear and tear. Inspect the 12-lead cable as described in the Operator's Checklist, and test it as described in Patient ECG Cable Check (on page 204).

Additional periodic preventive maintenance and testing—such as electrical safety tests, performance inspection, and required calibration—should be performed regularly by qualified service technicians. For detailed maintenance recommendations for each feature, see the *LIFEPAK 15 Monitor/Defibrillator Service Manual*.

General Maintenance and Testing

Table 36 Recommended Maintenance Schedule for Clinical Personnel

| OPERATION | DAILY | AFTER USE | AS REQUIRED | 6 MONTHS | 12 MONTHS |
|--|-------|--------------|----------------|-------------|--------------|
| Complete Operator's Checklist. Includes QUIK-COMBO therapy cable check and Standard (hard) paddles check | Х | | | | |
| Inspect defibrillator | Χ | Χ | | | |
| Check that all necessary supplies and accessories are present (for example, fully charged batteries, gel, electrodes, ECG paper, etc.) | Х | Х | Х | | |
| Function Checks: | | | | | |
| Patient ECG Cable Check (on page 204) | | | | Х | |
| Standard Paddles Synchronized Cardioversion Check (on page 205) | | | | Х | |
| Therapy Cable Monitoring and Synchronized Cardioversion Check (on page 206) | | | | Х | |
| Therapy Cable Pacing Check (on page 208) | | | | Х | |
| Clean defibrillator | | Х | Х | | |
| Preventive Maintenance and Testing | | | | | Х |

Self-Tests

Each time you turn on the LIFEPAK 15 monitor/defibrillator, it performs internal self-tests to check that internal electrical components and circuitry work properly. The defibrillator stores the results of all user-initiated self-tests in a test log.

When the defibrillator is on and a problem is detected that requires immediate service, such as a malfunctioning charging circuit, the Service LED is illuminated.

For more information, see General Troubleshooting Tips (on page 214).

Auto Tests

The defibrillator performs an automatic self-test daily at 03:00 (3:00 A.M.), if not in use. During the automatic self-test, the defibrillator turns itself on (**ON** LED illuminates) briefly and completes the following tasks:

- Performs a self-test
- Stores the self-test results in the test log
- Prints the self-test results

- Transmits the self-test results if the TRANSMIT RESULTS option is enabled. (Transmission may take up to 4 minutes.)
- · Turns itself off

If the defibrillator detects a problem during an auto test, it annotates the fault condition on the printed test report.

For more information about enabling the **TRANSMIT RESULTS** option, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* guide provided with your device.

The automatic self-test is not performed if the defibrillator is already turned on at 03:00 or if power is not available. If the defibrillator is manually turned on while a self-test is in progress, the self-test is halted and the defibrillator turns on normally.

For more information, see General Troubleshooting Tips (on page 214).

User Tests

The User Test is a functional test of the LIFEPAK 15 monitor/defibrillator. The User Test should be performed only as a test and not while using the defibrillator during patient care. Perform the User Test as a part of completing the daily Operator's Checklist.

Note: The defibrillator must be in Manual mode to perform the User Test.

To perform a User Test separate from completing the Operator's Checklist:

- 1. Press **ON** to turn on the LIFEPAK 15 monitor/defibrillator.
- 2. Press **OPTIONS**. The Options menu appears.
- 3. Select **USER TEST**. The defibrillator performs the following tasks:
 - · Self-tests to check the device.
 - Charges to 10 joules and discharges internally (this energy is not accessible at the therapy connector).
 - Prints a Pass/Fail report.

If the LIFEPAK 15 monitor/defibrillator detects a failure during the User Test, the Service LED is illuminated and the printed report indicates that the test failed. Remove the defibrillator from use and contact a qualified service technician.

If you must interrupt the User Test, turn the power off and then on again. The test stops and the defibrillator operates normally. A Pass/Fail report does not print.

Note: During the User Test, all front panel controls (except **ON**) and standard paddle controls are disabled. Routinely testing the defibrillator consumes battery power; maintain all batteries as described in Battery Warnings (on page 209).

Note: The last 40 User and Auto Test results are transmitted with all reports to the CODE-STAT Suite data management system.

Note: It is important to understand defibrillator operation. For suggested procedures to help keep personnel acquainted with normal defibrillator operation, see the function checks that are provided in this chapter. The function checks used may vary according to your local protocols.

General Maintenance and Testing

To test the defibrillator by performing the function checks, you need a simulator. To troubleshoot device performance, see General Troubleshooting Tips (on page 214).

Standard (Hard) Paddles Check

Perform the standard paddles check as a part of completing the daily Operator's Checklist that is provided in the back of this manual.

Function Checks

The following function checks are provided to help personnel keep acquainted with normal operating procedures and to troubleshoot LIFEPAK 15 monitor/defibrillator performance.

Note: If your organization downloads device electronic patient records for post-event review, consider entering "TEST" as the patient's name to distinguish simulator function tests from actual patient uses.

Patient ECG Cable Check

Equipment Needed:

- LIFEPAK 15 monitor/defibrillator
- Fully charged batteries or power adapter connected to a reliable power source
- Patient ECG cable (3-lead, 12-lead, or 5-wire)
- 3-lead or 12-lead simulator

To check the patient ECG cable:

- 1. Press ON.
- 2. Connect the ECG cable to the defibrillator.
- 3. Connect all cable leads to the simulator.
- 4. Turn on the simulator and select a rhythm.
- 5. Confirm that Lead II is selected.
- 6. After a few seconds, confirm that the screen displays a rhythm and that no **LEADS OFF** or **SERVICE** message appears.
- 7. For 12-lead cable, press **12-LEAD** and wait for printout. Confirm that a rhythm prints for each lead.

Standard Paddles Synchronized Cardioversion Check

WARNING

Shock Hazard

The defibrillator delivers up to 360 joules of electrical energy. Unless discharged properly as described in this test, this electrical energy may cause serious personal injury or death. Do not attempt to perform this test unless you are qualified by training and experience and are thoroughly familiar with these operating instructions.

Equipment Needed:

- LIFEPAK 15 monitor/defibrillator
- Standard paddles
- Defibrillator checker
- · Patient ECG cable
- · 3-lead or 12-lead patient simulator
- Fully charged batteries or power adapter connected to a reliable power source

To check standard paddles synchronized cardioversion:

- Press ON.
- 2. Connect the ECG cable to the monitor and to the patient simulator.
- 3. Turn on the simulator and select any rhythm except asystole or ventricular fibrillation.
- 4. Select Lead II.
- Press SYNC. Confirm that the SYNC LED lights. Adjust ECG size until the sense markers appear on the QRS complexes. Confirm that the SYNC LED blinks off with each detected QRS complex and that the heart rate is displayed.
- 6. Select 100 JOULES.
- 7. Press **CHARGE** and confirm that the tone indicating full charge sounds within 10 seconds or less.
- 8. Remove the standard paddles from the paddle wells and place the standard paddles on the defibrillator checker plates.

Note: This test is not intended to be performed with the paddles in the wells. Discharging 100 joules in the paddle wells may damage the defibrillator.

- 9. Press the APEX **₹** (shock) button, confirm that the defibrillator does not discharge, and then release the button.
- 10. Press the **STERNUM ►** (shock) button, confirm that the defibrillator does not discharge, and then release the button.
- 11.Press PRINT.

WARNING

Possible Paddle Damage and Patient Burns

Press paddles firmly onto the defibrillator checker plates when discharging to prevent arcing and formation of pits on paddle surfaces. Pitted or damaged paddles may cause patient skin burns during defibrillation.

- 12. Apply firm pressure with both paddles on the defibrillator checker paddle plates, and simultaneously press and hold both

 ✓ (shock) buttons while observing the screen.
- 13. Confirm that the defibrillator discharges on the next sensed QRS complex.
- 14. Press **PRINT** again to stop the printer.
- 15. Confirm that the defibrillator returns to Asynchronous mode (sense markers are no longer displayed and **SYNC** LED is off).

Note: Defibrillator may be set up to remain in Sync mode after discharge.

- 16. Confirm that the printer annotates the time, date, Sync On, sense markers prior to energy delivered, energy selected, no sense markers after Shock 1, and Sync Off on the ECG strip.
- 17. Turn off the defibrillator.

Note: If a **CONNECT CABLE, PADDLES LEADS OFF**, or any other warning message appears, replace the paddle assembly with a new paddle assembly and repeat the test. If the problem cannot be corrected, remove the device from active use and contact a qualified representative.

Therapy Cable Monitoring and Synchronized Cardioversion Check

CAUTION

Possible Simulator Damage

Do not discharge more than 30 shocks within an hour, or 10 shocks within a five-minute period, or pace continually into Physio-Control patient simulators. Simulators may overheat.

Equipment Needed:

- LIFEPAK 15 monitor/defibrillator
- QUIK-COMBO therapy cable
- Patient ECG cable
- 3-lead or 12-lead patient simulator with QUIK-COMBO connector
- Fully charged batteries or power adapter connected to a reliable power source

To check therapy cable monitoring and synchronized cardioversion:

- 1. Press ON.
- Connect the ECG cable to the defibrillator and to the simulator.
- 3. Connect the therapy cable to the simulator.
- 4. Turn on the simulator and select any rhythm except asystole or ventricular fibrillation.
- Select PADDLES lead.
- 6. Confirm that the screen displays an ECG and that the **PADDLES LEADS OFF** message does not appear.

Note: If the screen displays dashed lines, artifact (irregular noise signals), or any warning message, replace the therapy cable and repeat the test. If the problem cannot be corrected, remove the defibrillator from active use and contact a qualified service representative.

- 7. Select Lead II.
- Press SYNC. Confirm that the SYNC LED lights and the Sync mode message appears.
 Adjust ECG size until sense markers appear on the QRS complexes. Confirm that the
 SYNC LED blinks off with each detected QRS complex and that the heart rate is
 displayed.
- Select 50 JOULES.
- 10. Press CHARGE.
- 11. Press PRINT.



Shock Hazard

During defibrillation checks, the discharged energy passes through the cable connectors. Securely attach cable connectors to the simulator.

- 12. After the tone sounds indicating full charge, press and hold ₹ (shock) while observing the Home Screen.
- 13. Confirm that the defibrillator discharges on the next sensed QRS complex.
- 14. Press **PRINT** again to stop the printer.
- 15. Confirm that the defibrillator returns to Asynchronous mode (sense markers are no longer displayed and **SYNC** LED is off).

Note: Defibrillator may be set up to remain in Sync mode after discharge.

- 16. Select **PADDLES** lead.
- 17. Disconnect the therapy cable from the simulator. Confirm that the **PADDLES LEADS OFF** message appears and that an audible tone occurs.
- 18. Confirm that the printer annotates the time, date, Sync On, sense markers prior to energy delivered, energy selected, no sense markers after Shock 1, and Sync Off on the ECG strip.
- 19. Turn off the defibrillator.

Therapy Cable Pacing Check

Equipment Needed:

- LIFEPAK 15 monitor/defibrillator
- QUIK-COMBO therapy cable
- · Patient ECG cable
- 3-lead or 12-lead patient simulator with QUIK-COMBO connector
- Fully charged batteries or power adapter connected to a reliable power source

To check therapy cable pacing:

- 1. Press ON.
- 2. Connect the QUIK-COMBO therapy cable to the simulator.
- 3. Turn on the simulator and select BRADY.
- 4. Connect the ECG cable to the defibrillator and to the simulator.
- 5. Select Lead II.
- Press PACER.
- Confirm that sense markers appear on each QRS complex. If sense markers do not appear, or appear elsewhere on the ECG, press the SPEED DIAL on waveform Channel 1 and adjust ECG size from the menu.
- 8. Confirm that the RATE menu appears.
- Press CURRENT and increase the current to 80 mA.
- 10. Observe the screen for captured complexes. Confirm the **PACER** LED flashes with each delivered pacing pulse.
- 11. Disconnect the QUIK-COMBO therapy cable from the simulator. Confirm that the pacemaker stops pacing, the **CONNECT ELECTRODES** message appears, and an audible alarm sounds.
- 12. Reconnect the QUIK-COMBO therapy cable to the simulator. Confirm that the audible alarm stops, the **PACING STOPPED** message is displayed, and current is 0 mA.
- 13. Wait approximately 30 seconds and confirm that an audible alarm occurs.
- 14. Increase current to 80 mA. Confirm that audible alarm stops.
- 15. Press **CHARGE**. Confirm that the **PACER** LED goes off and that heart rate and available energy are displayed.

Battery Maintenance

This section provides information about the Physio-Control Lithium-ion batteries that are specifically designed for use in the LIFEPAK 15 monitor/defibrillator. Lithium-ion batteries are low maintenance and require no scheduled cycling to prolong battery life.

IMPORTANT! The LIFEPAK 15 monitor/defibrillator Lithium-ion batteries, battery chargers, power adapters, and power cords are not interchangeable with batteries, battery chargers, power adapters, and power cords that are used in other LIFEPAK defibrillators.

Battery Warnings

WARNING

Possible Fire, Explosion, and Burns

Do not disassemble, puncture, crush, heat above 100°C (212°F), or incinerate the battery.

WARNING

Possible Loss of Power and Delay of Therapy During Patient Care

Using an improperly maintained battery to power the defibrillator may cause power failure without warning. Use the appropriate Physio-Control battery charger to charge batteries.

WARNING

Possible Loss of Power During Patient Care

Physio-Control has no information regarding the performance or effectiveness of its LIFEPAK monitor/defibrillators if other manufacturers' batteries, battery chargers, or power adapters are used. Using other manufacturers' batteries, battery chargers, or power adapters may cause the device to perform improperly and invalidate the safety agency certifications. Use only Physio-Control LIFEPAK 15 monitor/defibrillator batteries (PN 3206735) and the appropriate Physio-Control LIFEPAK 15 monitor/defibrillator battery charger or power adapter.

WARNING

Possible Loss of Power During Patient Care

Battery pins in the defibrillator may be damaged if batteries are dropped or forced into battery wells. Inspect pins routinely for signs of damage. Keep batteries installed at all times except when device is removed from service for storage.

CAUTION

Possible Equipment Damage

When storing the LIFEPAK 15 monitor/defibrillator for an extended period of time, the battery should be removed from the device.

Receiving New Batteries

New batteries do not arrive fully charged. Charge each new battery before use. Batteries may be charged using any of the following devices:

- Station Lithium-ion battery charger for use with the LIFEPAK 15 monitor/defibrillator
- Mobile Lithium-ion battery charger for use with the LIFEPAK 15 monitor/defibrillator
- REDI-CHARGE battery charger
- AC power adapter for use with the LIFEPAK 15 monitor/defibrillator
- DC power adapter for use with the LIFEPAK 15 monitor/defibrillator

Storing Batteries

Li-ion batteries self-discharge during storage.

If you store the battery:

- Do not remove the Charge Before Use label to indicate that the battery has not yet been charged.
- Store batteries at temperatures between 20° to 25°C (68° to 77°F).
- Charge the battery fully within one year of when you receive it. Fully recharge the battery once per year thereafter.

WARNING

Possible Loss of Power During Patient Care

Stored batteries lose charge. Failure to charge a stored battery before use may cause device power failure without warning. Always charge a stored battery before placing it in active use.

Charging Batteries

- Charge batteries before use. Batteries may be charged in a battery charger, or in the LIFEPAK 15 monitor/defibrillator if it is connected to an auxiliary power source using a LIFEPAK 15 monitor/defibrillator power adapter.
- Inspect batteries for damage or leakage. If battery is damaged or leaking, recycle the battery and obtain a new battery.
- Remove the Charge Before Use label from new batteries before placing batteries in the charger or in the LIFEPAK 15 monitor/defibrillator.
- The battery fuel gauge does not function until the battery is charged. For more information about the fuel gauge, see Batteries (on page 32).
- For more information about charging batteries, refer to either the *Instructions for Use* provided with your battery charger, or the Power Adapter (on page 189) chapter if using the power adapter.

Replacing Batteries

Physio-Control recommends that batteries be replaced approximately every two years. Properly maintained batteries may last longer. A battery has reached the end of useful life if *one or more* of the following circumstances occur:

- Physical damage occurs to the battery case, for example, cracks or a broken clip.
- The battery is leaking.
- The battery charger indicates **FAULT**.
- The battery fuel gauge indicates two or fewer LEDs (bars) after the battery completes a charge cycle.

Dispose of used batteries promptly. Keep batteries away from children.

Recycling Batteries

To promote awareness of battery recycling, Physio-Control batteries are marked with one of these symbols:









When a battery has reached the end of its useful life, recycle the battery as described below.

Battery Recycling in the USA

Recycle batteries by participating with Physio-Control in a national recycling program. Contact your Physio-Control representative to obtain shipping instructions and shipping containers. Do not return your batteries to the Physio-Control offices in Redmond, Washington, unless instructed to do so.

Battery Recycling Outside the USA

Recycle batteries according to national and local regulations. Contact your local Physio-Control representative for assistance.

Cleaning the Device

CAUTION

Possible Equipment Damage

Do not clean any part of this device or its accessories with bleach, bleach dilution, or phenolic compounds. Do not use abrasive or flammable cleaning agents. Do not attempt to sterilize this device or any accessories unless otherwise specified in accessory operating instructions.

Clean the LIFEPAK 15 monitor/defibrillator, therapy and ECG cables, and batteries with a damp sponge or cloth. Use only the cleaning agents listed below:

- · Quaternary ammonium compounds
- · Isopropyl alcohol
- Peracetic (peroxide) acid solutions

Note: Carefully clean the connector ports. Do not allow cleaning fluids to penetrate the exterior surfaces of the device.

Clean the carrying case accessory as follows and as described on its instruction tag:

 Hand wash using mild soap or detergent and water. A scrub brush may be useful for heavily soiled spots. Cleaners such as Formula 409[®] are helpful for grease, oil, and other tough stains.

For information about cleaning the reusable monitoring sensors and cables, see the individual monitoring section.

Storing the Device

To take the LIFEPAK 15 monitor/defibrillator out of service and store it for an extended period of time, follow these guidelines:

- Remove the batteries.
- Store the defibrillator and batteries at room temperature.

For more information about storage and operating specifications, see Environmental Specifications (on page 235).

To return the LIFEPAK 15 monitor/defibrillator to service, perform the following tasks:

- Complete the tasks listed in the Operator's Checklist located at the end of this manual. If the Operator's Checklist can not be located, a copy is available at www.physio-control.com.
- Consider having the device serviced by a qualified service technician.

Loading Paper

Check the amount of paper in the printer as part of the daily check according to the Operator's Checklist provided in the back of this manual.

CAUTION

Possible Printer Malfunction

Using other manufacturers' printer paper may cause the printer to function improperly or damage the print head. Use only Physio-Control printer paper.

The printer is equipped with an out-of-paper sensor to protect the printer printhead. The sensor automatically turns off the printer if paper runs out or the printer door is open.

To load paper:

- 1. Lift the printer door latch to release the door (see Figure, Loading Paper).
- 2. Pull out the printer door.
- 3. Remove the empty paper spool, if present.
- 4. Insert a new paper roll with the graph side facing up. Make sure the end of the paper extends outward so it is exposed when the printer door is closed.
- 5. Close the printer door and press down on the latch until the door clicks shut.

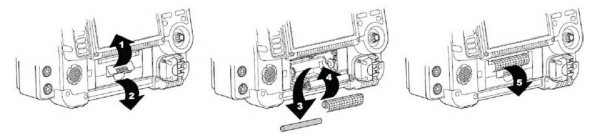


Figure 53 Loading Paper

General Troubleshooting Tips

If a problem is detected with the LIFEPAK 15 monitor/defibrillator during operation or testing, refer to the following troubleshooting tips. If the problem cannot be corrected, remove the LIFEPAK 15 monitor/defibrillator from active use and contact a qualified service technician for service and repair.

Table 37 General Troubleshooting Tips

| No power when L monitor/defibrillator is turned ON | POSSIBLE CAUSE Low battery voltage | Replace with fully charged, properly maintained battery. |
|--|---|--|
| monitor/defibrillator is turned ON | · · | |
| | - | |
| | Battery connector pin loose, covered with foreign substance, or damaged | Remove battery and inspect pins. Clean if foreign substance present. Contact a qualified service technician to replace if bent, cracked, or loose. |
| C | Power adapter not properly connected to auxiliary power source | Check that power adapter is properly connected to auxiliary power. |
| C | Power adapter not properly connected to monitor/defibrillator | Check that power adapter is properly connected to monitor/defibrillator. |
| | Defective power adapter or cables | Replace with working power adapter and cables. |
| | Defective battery | Remove battery from service and replace with working battery. |
| ON LED illuminated, but screen is blank and device does not operate | Device boot up has failed | Press and hold ON until LED turns off (~5 seconds). Then press ON to turn device back on. |
| | | If device does not turn off, remove both batteries and disconnect device from power adapter, if applicable. Then reinsert batteries, reconnect power adapter, and press ON to turn device back on. |
| not illuminated c | Power adapter not properly connected to auxiliary power source | Check that power adapter is properly connected to auxiliary power. |
| C | Power adapter not properly connected to monitor/defibrillator | Check that power adapter is properly connected to monitor/defibrillator. |
| | Defective power adapter or cables | Replace with working power adapter and cables. |
| on monitor/defibrillator not dilluminated | Power adapter not properly connected to auxiliary power source or monitor/defibrillator | Check that power adapter is connected properly. |
| | Battery not properly inserted in battery well | Check that battery is properly inserted in battery well. |

Chapter 10 | Maintaining the Equipment

| Unable to charge battery with power adapter because battery charge level is too low. Contect of REDI-CHARGE battery charge rif available. Replace battery charge rif available. Replace battery. | ODOEDWATION | DOCCIDI E CALICE | |
|--|-----------------------|----------------------------|---|
| with power adapter because battery charge level is too low. No batteries installed Pefective battery Defective battery Inrecognized battery Unrecognized battery Incompatible power adapter connected to the monitor/defibrillator. Defective power adapter or cables Monitor/defibrillator unable to recognize installed battery Defective power adapter or cables Defective power adapter or cables | OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
| No batteries installed No batteries installed No batteries installed No battery | | with power adapter because | REDI-CHARGE battery charger if |
| Defective battery . Remove battery from service and replace with working battery. | | low. | Replace battery. |
| Unrecognized battery Only use battery that is approved for use with the LIFEPAK 15 monitor/defibrillator. | | No batteries installed | Install at least one battery. |
| Incompatible power adapter connected to the monitor/defibrillator | | Defective battery | |
| connected to the monitor/defibrillator Defective power adapter or cables Monitor/defibrillator unable to recognize installed battery Defective battery Defective power adapter appears Defective battery Defective power adapter and cables. Contact qualified service personnel. Remove battery from service and replace with working battery. Replace with working power adapter and cables. Contact qualified service personnel. Charge battery in Station-Mobile or REDI-CHARGE battery charger. Replace battery. Press ON immediately to turn device back on. Replace battery immediately. Press ON to turn device back on. Replace battery immediately. Press ON to turn device back on. Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough • Press and hold ON for at least two | | Unrecognized battery | use with the |
| cables Monitor/defibrillator unable to recognize installed battery Defective battery Defective battery Defective power adapter Device unable to charge battery or batteries Fuel gauge on battery does not illuminate Extremely depleted battery Faulty battery Paulty battery Press ON immediately to turn device back on. RF equipment too close to defibrillator Distances (on page 266). Press ON to turn device back on. Cellular equipment too close to installed battery Cellular equipment too close to to installed battery LIFENET Gateway (modem) too close to installed battery Device won't turn off ON not pressed long enough CANNOT CHARGE Press and working battery are replace with working power adapter and cables. Contact qualified service and replace with working battery. Pcharge battery im Station-Mobile or REDI-CHARGE battery in Station-Mobile or REDI-CHARGE battery. Press ON immediately to turn device back on. Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. If device does not turn on, replace battery. Press ON to turn device back on. If device does not turn on, replace battery. Press ON to turn device back on. | | connected to the | approved for use with the |
| CANNOT CHARGE BATTERY message appears Defective battery Defective power adapter Device unable to charge battery or batteries Fuel gauge on battery does not illuminate Extremely depleted battery Faulty battery Press ON immediately to turn device back on. Device turns off unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpectedly Device turns off Unexpected battery Press ON to turn device back on. Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. Move cellular equipment away from installed battery. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | | |
| BATTERY message appears Defective power adapter Device unable to charge battery or batteries Fuel gauge on battery does in tilluminate Extremely depleted battery Faulty battery Pavice turns off unexpectedly Press ON inturn device back on. RF equipment too close to defibrillator Cellular equipment too close to installed battery Cellular equipment too close to installed battery LIFENET Gateway (modem) too close to installed battery. Device won't turn off ON not pressed long enough Press ON tourn device back on. Replace with working power adapter and cables. Contact qualified service personnel. Charge battery in Station-Mobile or REDI-CHARGE battery charger. Press ON immediately to turn device back on. Replace battery immediately. Press ON to turn device back on. Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. Move cellular equipment away from installed battery. Press ON to turn device back on. If device does not turn on, replace battery. Press ON to turn device back on. If device does not turn on, replace battery. Press ON to turn device back on. | | | Contact qualified service personnel. |
| Device unable to charge battery or batteries Fuel gauge on battery does not illuminate Extremely depleted battery Faulty battery Faulty battery Peress ON immediately to turn device back on. Faculty power Low battery power RF equipment too close to defibrillator Cellular equipment too close to installed battery Cellular equipment too close to installed battery LIFENET Gateway (modern) too close to installed battery Device won't turn off ON not pressed long enough Contact qualified service personnel. Charge battery in Station-Mobile or REDI-CHARGE battery in Station-Mobile or REDI-CHARGE battery in Station-Mobile or REDI-CHARGE battery. Press ON immediately to turn device back on. Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. Move cellular equipment away from installed battery. Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | BATTERY message | Defective battery | |
| Fuel gauge on battery does not illuminate Extremely depleted battery not illuminate Extremely depleted battery Faulty battery Faulty battery Press ON immediately to turn device back on. Low battery power Low battery power Press ON to turn device back on. RF equipment too close to defibrillator Press ON to turn device back on. Cellular equipment too close to installed battery Cellular equipment too close to installed battery LIFENET Gateway (modem) too close to installed battery Device won't turn off ON not pressed long enough Press and hold ON for at least two | | Defective power adapter | |
| not illuminate Faulty battery Press ON immediately to turn device back on. Low battery power Faulty power Faulty power Faulty battery Press ON immediately to turn device back on. Replace battery immediately. Press ON to turn device back on. Separate RF equipment from defibrillator Distances (on page 266). Press ON to turn device back on. Cellular equipment too close to installed battery Fress ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | | Contact qualified service personnel. |
| Device turns off unexpectedly High power draw Low battery power RF equipment too close to defibrillator RF equipment too close to defibrillator Cellular equipment too close to installed battery LIFENET Gateway (modem) too close to installed battery LIFENET Gateway (modem) too close to installed battery Device won't turn off ON not pressed long enough Press ON immediately to turn device back on. Replace battery immediately. Replace battery immediately. Press ON to turn device back on. Move cellular equipment away from installed battery. Press ON to turn device back on. If device does not turn on, replace battery. Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | Extremely depleted battery | |
| Low battery power Evaluate too close to defibrillator | | Faulty battery | Replace battery. |
| Press ON to turn device back on. RF equipment too close to defibrillator Separate RF equipment from defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. Cellular equipment too close to installed battery Move cellular equipment away from installed battery. Press ON to turn device back on. If device does not turn on, replace battery. Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | High power draw | _ |
| defibrillator defibrillator. See Separation Distances (on page 266). Press ON to turn device back on. Cellular equipment too close to installed battery Press ON to turn device back on. Hove cellular equipment away from installed battery. Press ON to turn device back on. If device does not turn on, replace battery. Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | Low battery power | - |
| Cellular equipment too close to installed battery • Move cellular equipment away from installed battery. • Press ON to turn device back on. • If device does not turn on, replace battery. LIFENET Gateway (modem) too close to installed battery • Store modem in side pouch of defibrillator. Do not store modem in back pouch. • Press ON to turn device back on. • If device does not turn on, replace battery. Device won't turn off ON not pressed long enough • Press and hold ON for at least two | | | defibrillator. See Separation Distances (on page 266). |
| to installed battery Press ON to turn device back on. If device does not turn on, replace battery. LIFENET Gateway (modem) too close to installed battery Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | | Press ON to turn device back on. |
| LIFENET Gateway (modem) too close to installed battery Store modem in side pouch of defibrillator. Do not store modem in back pouch. Press ON to turn device back on. If device does not turn on, replace battery. Device won't turn off ON not pressed long enough Press and hold ON for at least two | | | |
| LIFENET Gateway (modem) too close to installed battery • Store modem in side pouch of defibrillator. Do not store modem in back pouch. • Press ON to turn device back on. • If device does not turn on, replace battery. Device won't turn off • Press and hold ON for at least two | | | • Press ON to turn device back on. |
| too close to installed battery defibrillator. Do not store modem in back pouch. • Press ON to turn device back on. • If device does not turn on, replace battery. Device won't turn off ON not pressed long enough • Press and hold ON for at least two | | | |
| • If device does not turn on, replace battery. Device won't turn off ON not pressed long enough • Press and hold ON for at least two | | | defibrillator. Do not store modem in |
| Device won't turn off ON not pressed long enough Press and hold ON for at least two | | | • Press ON to turn device back on. |
| Device won't turn off ON not pressed long enough • Press and hold ON for at least two | | | |
| | Device won't turn off | | Press and hold ON for at least two |

General Troubleshooting Tips

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---|--|--|
| Monitor/defibrillator operates, but screen is blank | Operating temperature is too low or too high | |
| | Screen not operating properly | Print ECG strip to assess rhythm and other active vital signs. |
| | | Press ANALYZE and use AED mode, if necessary. |
| | | Contact qualified service technician. |
| Monitor/defibrillator operates, but screen not | Screen in direct sunlight | Change screen from color to black and white. |
| readable | | Reposition or shield device. |
| | | Print ECG strip to assess rhythm and other active vital signs. |
| | | Press ANALYZE and use AED mode, if necessary. |
| CHECK PRINTER message | Printer paper jams, slips, or | Reinstall paper. |
| appears | misfeeds | If problem persists, contact qualified service technician. |
| | Printer is out of paper | Add new paper. |
| Service LED illuminates | Device self-test circuitry detects service condition | Continue to use defibrillator or pacemaker, if needed. |
| | | Turn device off and then on again. Note that this creates a new "patient." If Service LED does not clear, remove device from active use. |
| | | Report occurrence of Service LED to qualified service personnel. |
| | | Obtain another defibrillator, if necessary. |
| ECG monitoring problems | | See Troubleshooting Tips (on page 55). |
| Problems with AED operation | | See Troubleshooting Tips (on page 130). |
| Problems with defibrillation/synchronized cardioversion | | See Troubleshooting Tips (on page 139). |
| Problems with pacing | | See Troubleshooting Tips (on page 145). |
| Displayed time is incorrect | Time is incorrectly set | Change the time setting. See Options (on page 41). |
| Date printed on report is incorrect | Date is incorrectly set | Change the date setting. See Options (on page 41). |
| Displayed messages are faint or flicker | Low battery power Out of temperature range | Replace the battery immediately.Connect to auxiliary power using approved power adapter. |
| Low speaker volume | Moisture in speaker grill holes | Wipe moisture from speaker grill and allow device to dry. |
| | | |

Chapter 10 | Maintaining the Equipment

| OBSERVATION | POSSIBLE CAUSE | CORRECTIVE ACTION |
|---------------------------------|--|--|
| MAINTENANCE DUE message appears | Maintenance prompt is set to display at a selected | Continue to use device, if needed.Contact service personnel to |
| | interval in Service mode | perform routine maintenance. Contact Physio-Control Technical Support for instructions on how to reset or turn off this prompt. |

Service and Repair

WARNING Shock Hazard Do not disassemble the defibrillator. It contains no operator serviceable components and dangerous high voltages may be present. Contact a qualified service technician for repair. WARNING Ineffective Energy Delivery Hazard Service mode is for authorized personnel only. Improper use of Service mode may inappropriately alter the device's configuration and may change energy output levels. Contact qualified service technician for assistance or information about device configuration.

If the LIFEPAK 15 monitor/defibrillator requires service as indicated by testing, troubleshooting, or a service message, contact a qualified service technician. In the USA, call Physio-Control Technical Support at 1.800.442.1142.

When calling Physio-Control to request service, identify the model and serial number and describe the observation. If the device must be shipped to a service center or the factory, pack the device in the original shipping container, if possible, or in protective packing to prevent shipping damage.

The LIFEPAK 15 Monitor/Defibrillator Service Manual provides detailed technical information to support service and repair by a qualified service technician.

Product Recycling Information

Recycle the device at the end of its useful life.

Recycling Assistance

The device should be recycled according to national and local regulations. Contact your local Physio-Control representative for assistance or refer to www.physio-control.com/recycling.

Preparation

The device should be clean and contaminant-free prior to being recycled.

Recycling of Disposable Electrodes

After using disposable electrodes, follow your local clinical procedures for recycling.

Packaging

Packaging should be recycled according to national and local regulations.

Warranty

To obtain a detailed warranty statement, contact your local Physio-Control representative or go to www.physio-control.com.

Using defibrillation electrodes, adapter devices, or other parts and supplies from sources other than Physio-Control is not recommended. Physio-Control has no information regarding the performance or effectiveness of its LIFEPAK defibrillators if they are used in conjunction with defibrillation electrodes or other parts and supplies from other sources. If device failure is attributable to defibrillation electrodes or other parts or supplies not manufactured by Physio-Control, this may void the warranty.

Accessories

The following table lists accessories that are available for the LIFEPAK 15 monitor/defibrillator. To order, contact your Physio-Control representative. In the USA, call Customer Support at 1.800.442.1142, option 2.

Note: The LIFEPAK 15 monitor/defibrillator and its accessories that are intended for direct or casual contact with the patient are not made with natural rubber latex.

Table 38 Accessories for the LIFEPAK 15 Monitor/Defibrillator

| Table 38 Accessories for | or the LIFEPAK 15 Monitor/Defibrillator |
|----------------------------|---|
| CATEGORY | RELATED ACCESSORY |
| Power | Lithium-ion battery |
| | Station Lithium-ion Battery Charger |
| | Mobile Lithium-ion Battery Charger |
| | REDI-CHARGE Battery Charger |
| | AC Power Adapter for use with the LIFEPAK 15 monitor/defibrillator |
| | DC Power Adapter for use with the LIFEPAK 15 monitor/defibrillator |
| | Power adapter output extension cable |
| | Power adapter attachment kit |
| Therapy | QUIK-COMBO pacing/defibrillation/ECG electrodes |
| | QUIK-COMBO RTS pacing/defibrillation/ECG electrodes |
| | QUIK-COMBO RTS Pediatric pacing/defibrillation/ECG electrodes |
| | QUIK-COMBO pacing/defibrillation/ECG electrodes with REDI-PAK |
| | preconnect system |
| | QUIK-COMBO Therapy cable |
| | Standard paddles |
| | Pediatric paddles |
| | Internal paddles |
| | Internal paddles adapter cable |
| Monitoring: | |
| ECG | Cleartrace™ ECG electrodes (Conmed) |
| | 3-lead ECG cable |
| | 5-wire ECG cable |
| | 12-lead ECG cable (includes main 4-wire cable and precordial lead attachment) |
| SpO ₂ - Masimo | RC-4 patient cable (4 ft) |
| | RC-12 patient cable (12 ft) |
| | Red LNC patient cable (4, 10, 14 ft) |
| | Patient extension cables Red™ LNOP® and LNCS™ |
| | Reusable LNCS and M-LNCS sensors |
| | Disposable LNCS and M-LNCS sensors |
| SpO ₂ – Nellcor | Masimo Red™ MNC patient cable (for use with Nellcor sensors) |
| | Reusable Oximax DS-100A sensor |
| | Dura-Y™ multisite sensor |
| | Oxiband reusable sensor, Adult/Neonatal |

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Accessories

| CATEGORY | RELATED ACCESSORY |
|-----------------------------|---|
| SpCO and SpMet - | Rainbow patient extension cables |
| Masimo | Rainbow reusable sensors |
| | Rainbow disposable sensors |
| | Rainbow light shields |
| NIBP | NIBP reusable blood pressure cuffs (Statcorp Medical) |
| | NIBP disposable blood pressure cuffs (Statcorp Medical) |
| | NIBP hoses |
| EtCO ₂ – Oridion | EtCO ₂ FilterLine sets |
| | EtCO₂ Smart CapnoLine® lines |
| Temperature | Measurement Specialties disposable temperature probes: 4491 Esophageal/Rectal, 4499HD Skin High Dielectric, 4464 Foley 14Fr, 4466 Foley 16Fr, 4468 Foley 18Fr |
| | Temperature probe adapter cable |
| Other accessories | Wireless modem/gateway |
| | Bed connector |
| | LIFEPAK monitor to PC cable (serial communication cable) |
| | PC-based configuration tool |
| | Test Load |
| | 3-Lead ECG patient simulator |
| | 12-Lead ECG patient simulator |
| | SIGNAGAL® electrode gel |
| | ECG recording paper, 100 mm wide |

Appendix A

Specifications and Performance Characteristics

This appendix contains the specifications and performance characteristics for the LIFEPAK 15 monitor/defibrillator and the LIFEPAK 15 monitor/defibrillator batteries. It also lists high and low alarm limits, alarm performance characteristics, and factory default settings.

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Specifications and Performance Characteristics

Table, LIFEPAK 15 Monitor/Defibrillator Specifications, lists the LIFEPAK 15 monitor/defibrillator specifications for the device.

Table, Battery Specifications (on page 236), lists the specifications for the LIFEPAK 15 monitor/defibrillator batteries.

Table, Alarm Limits (on page 237), lists the high and low limits for alarms when either the wide or narrow alarm setting is selected on the LIFEPAK 15 monitor/defibrillator.

Table, Alarm Performance Characteristics (on page 237), lists the alarm performance characteristics.

Table, Setup Options Factory Default Settings (on page 239), lists the factory default settings for the LIFEPAK 15 monitor/defibrillator setup options.

 Table 39 LIFEPAK 15 Monitor/Defibrillator Specifications

| CHARACTERISTIC All specifications are at 20° | DESCRIPTION C unless otherwise stated. |
|---|--|
| GENERAL | |
| Classification | Monitor/defibrillator—Battery powered and Class II (per IEC 60601-1) |
| | Applied parts—ECG, Internal Defibrillation, Invasive Pressure and Temperature have Type CF patient connections. External Defibrillation, CO2, SpO2, and NIBP have Type BF patient connections (per IEC 60601-1). |
| Modes | AED mode—for automated ECG analysis and a prompted treatment protocol for patients in cardiac arrest. Manual mode—for performing manual defibrillation, synchronized cardioversion, noninvasive pacing, and ECG and vital sign monitoring. Archive mode—for accessing stored patient information. Setup mode—for changing default settings of the operating functions. Service mode—for authorized personnel to perform diagnostic tests and calibrations. Demo mode—for simulated waveforms and trend graphs for demonstration purposes. |
| Self-test | When powered on, the device performs a self-test to check internal electrical components and circuitry. A service indicator is illuminated if an error is detected. The device also performs an auto test daily. Results are printed and stored in the device log. Auto test results can be transmitted. See the LIFEPAK 15 Monitor/Defibrillator Setup Options provided with your device for more information. |
| Continuous Patient Surveillance System (CPSS) | In Advisory Monitoring, CPSS monitors the patient ECG, via QUIK-COMBO® electrodes or Lead II, for a potentially shockable rhythm. |
| Voice Prompts | Manual mode: Used for selected prompts (selectable ON/OFF) AED mode: Used for entire AED protocol |

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Specifications and Performance Characteristics

| Analog ECG | à Output | Output: 1 volt/mV Frequency Response: 0. ECG and 1.3 to 23 Hz fo No internal pacemaker p included. Pace markers included. | or 1–30 Hz Monitor Fred oulse enhancements or | quency Response) detection markers |
|--------------|--|---|---|--|
| Notch Filter | | 50 or 60 Hz | | |
| POWER | | | | |
| Batteries | | Rechargeable Lithium-io | n battery | |
| | | Dual battery capability w | vith automatic switching | g |
| | | Low battery indication a indication and low batter | | |
| | | Replace battery indication indication, audio tones, a area for each battery. We switches to second batter condition, a voice prompt | and replace battery me hen replace battery is i ery. When both batterie | essage in the status ndicated, device auto- es reach replace battery |
| | | Input voltage range is between +8.8 and +12.6 Vdc | | Vdc |
| Battery Cap | acity | For two, new fully-charg | ed batteries, 20°C (68° | F): |
| | | Capacity to shutdown is | : | |
| Oper | rating Mode | Monitoring (minutes) | Pacing (minutes) | Defibrillation (360J discharges) |
| | Typical | 360 | 340 | 420 |
| | Minimum | 340 | 320 | 400 |
| | | Capacity after low batter | ry is: | |
| _ | Typical | 21 | 20 | 30 |
| | Minimum | 12 | 10 | 6 |
| AC Power A | dapter | AC-DC power adapter | | |
| | | Input power range is 100-240 Vac, 50/60 Hz, 1.4-0.6 A | | |
| | | Output voltage is 12 Vdc | | |
| | | Meets UL 60601-1 300 r installed on a center-tap | | |
| DC Power A | Adapter | DC-DC power adapter | | |
| | | Input power range is: Minimum: 11 Vdc, 15 Nominal: 13.8 Vdc, 1 Maximum: 17.6 Vdc, | 2.5 A | |
| | | Output voltage is 12 Vdc | ; | |
| | ce Behavior when using wer Adapter | Auxiliary power indicator to auxiliary power. Batte batteries are fully charge charged. | ry charging indicator ill | uminated when |
| | | Battery status indicators number is not highlighte and replace battery pron | d because battery is no | ot in use. Low battery |

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Appendix A | Specifications and Performance Characteristics

| PHYSICAL | |
|--|--|
| Weight | Basic monitor/defibrillator with new roll paper and two batteries installed: 7.9 kg (17.5 lb) |
| | Fully featured monitor/defibrillator with new roll paper and two batteries installed: 8.4 kg (18.5 lb) |
| | Lithium-ion battery: < 0.60 kg (1.3 lb) |
| | Accessory bags and shoulder strap: 1.77 kg (3.9 lb) |
| | Standard (hard) paddles: 0.95 kg (2.1 lb) |
| Height | 31.7 cm (12.5 in) |
| Width | 40.1 cm (15.8 in) |
| Depth | 23.1 cm (9.1 in) |
| DISPLAY | |
| Size (active viewing area) | 212 mm (8.4 in) diagonal; 171 mm (6.7 in) wide x 128 mm (5.0 in) high |
| Display Type | 640 dot x 480 dot color backlit LCD |
| | User selectable display mode (full color or SunVue™ high contrast) |
| | Displays a minimum of 5 seconds of ECG and alphanumerics for values, device instructions, or prompts |
| | Displays up to three waveforms |
| | Waveform display sweep speed: 25 mm/sec for ECG, SpO ₂ , IP, and 12.5 mm/sec for CO ₂ |
| DATA MANAGEMENT | |
| | nd stores patient data, events (including waveforms and annotations), and patient impedance records in internal memory. |
| The user can select and communication method | d print reports, and transfer the stored information via supported ds. |
| Report Types | Three format types of CODE SUMMARY™ critical event record: short, medium, and long |
| | 12-lead ECG with STEMI statements |
| | Continuous Waveform (transfer only) |
| | Trend Summary |
| | Vital Sign Summary |
| | Snapshot |
| Memory Capacity | Total capacity is 360 minutes of continuous ECG, 90 minutes of continuous data from all channels, or 400 single waveform events. Maximum memory capacity for a single patient includes up |
| | to 200 single waveform reports and 90 minutes of continuous ECG. |

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Specifications and Performance Characteristics

| COMMUNICATIONS | |
|---|--|
| complies with Part 15 of conditions: (1) this devic | transferring data records by wired or wireless connection. This device the FCC rules, and its operation is subject to the following two e may not cause harmful interference, and (2) this device must accept d, including interference that may cause undesired operation. |
| Serial Port | RS232 communication +12V available |
| Bluetooth® technology | Bluetooth technology provides short-range wireless communication with other Bluetooth-enabled devices. The Bluetooth transceiver complies with Bluetooth Class 1 frequency, power, and bandwidth requirements. |
| MONITOR | |
| ECG | ECG is monitored via several cable arrangements. A 3-wire cable is used for 3-lead ECG monitoring. A 5-wire cable is used for 7-lead ECG monitoring. A 10-wire cable is used for 12-lead ECG acquisition When the chest electrodes are removed, the 10-wire cable functions as a 4-wire cable. Standard paddles or QUIK-COMBO pacing/defibrillation/ECG electrodes are used for paddles lead monitoring. |
| Frequency Response | Monitor—0.5 to 40 Hz or 1 to 30 Hz Paddles—2.5 to 30 Hz 12-lead ECG diagnostic—0.05 to 150 Hz |
| Lead Selection | Leads I, II, III (3-wire ECG cable) |
| | Leads I, II, III, AVR, AVL, and AVF acquired simultaneously (4-wire ECG cable) |
| | Leads I, II, III, AVR, AVL, AVF, and C lead acquired simultaneously (5-wire ECG cable) |
| | Leads I, II, III, AVR, AVL, AVF, V1, V2, V3, V4, V5, and V6 acquired simultaneously (10-wire ECG cable) |
| ECG Size | 4, 3, 2.5, 2, 1.5, 1, 0.5, 0.25 cm/mV (fixed at 1 cm/mV for 12-lead) |
| Leads Off Sensing | The ECG leads off function uses AC current at 20 kHz for sensing leads off, the disposable defibrillation electrodes use AC current at 20 kHz for leads off, and the ECG leads use a noise cancelation signal which ranges from DC to approximately 5 kHz. The amplitude of these signals conforms to IEC 60601-1 Clause 8.7.3. |
| Heart Rate Display | 20–300 bpm digital display |
| | Accuracy: ±4% or ±3 bpm, whichever is greater |
| Recovery Time after Defibrillation | 10 seconds |
| Heart Rate Averaging Method | The heart rate average is formed by a weighted average of approximately 8 seconds duration. When the input rate is trending rapidly, the rate meter will track more quickly. Refer to heart rate response time disclosure. The display update interval is every heartbeat or every 2 seconds, whichever is shorter. |
| Heart Rate Step Response Time | 80 bpm to 120 bpm input step change: ≤ 10 seconds to indicate a minimum of 100 bpm. 80 bpm to 40 bpm input step change: ≤ 10 seconds to indicate a maximum of 60 bpm. |

Appendix A | Specifications and Performance Characteristics

| Heart Rate with Irregular Rhythm | The rate meter output can range from the heart rate associated with the shortest R-R interval to the heart rate associated with the longest R-R interval. When present, intermediate length R-R intervals are favored as the basis for the rate. When evaluated per IEC 60601-2-27, rates are as follows: • A1. Ventricular bigeminy: HR = 80 to 86 • A2. Slow alternating ventricular bigeminy: HR = 60 to 63 • A3. Rapid alternating ventricular bigeminy: HR = 123 to 124 • A4. Bidirectional systoles: HR = 97 to 99 |
|---------------------------------------|---|
| QRS Detection Range | Duration: 40 to 120 msec Amplitude: 0.5 to 5.0 mV Tall T-wave Rejection: T-waves that are 1 mV high are not detected b the monitor when the R-wave size is 1 mV and input rate is 80 ppm. |
| Common Mode Rejection (CMRR) | ECG Leads: 90 dB at 50/60 Hz |
| Pacemaker Pulse Rejection | Rejects pacemaker pulses having amplitudes from ±2 mV to ±700 mV and pulse widths from 0.1 ms to 2.0 ms with and without overshoot. Pacemaker pulse overshoot is defined as 2.5% to 25% of the pacemaker pulse amplitude not to exceed 2 mV. Refer to IEC 60601-2-27. Note: Does not apply when internal paddles are connected. |
| 0.0 /0.0 00 /0.0 Mad | Note: Does not apply when internal paddles are connected. |
| SpO ₂ /SpCO/SpMet | Masina e e e e e e e e e e e e e e e e e e e |
| Sensors | Masimo [®] sensors including Rainbow [®] sensors Nellcor [®] sensors when used with the Masimo Red™ MNC adapter |
| SpO ₂ | |
| Displayed Saturation Range | "<50" for levels below 50%; 50 to 100% |
| Saturation Accuracy is sp | pecified for range 70-100% (0-69% is not specified). |
| Adults/Pediatrics Accuracy (RMS)* | ±2% (during no motion conditions - Masimo) ±3% (during no motion conditions - Nellcor) ±3% (during motion conditions - Masimo) |
| Neonatal Accuracy (RMS)* | ±3% (during no motion conditions - Masimo) ±4% (during no motion conditions - Nellcor ±3% (during motion conditions - Masimo) |
| Dynamic signal strength | |
| Pulse tone as SpO ₂ pulsa | itions are detected |
| SpO ₂ Averaging Time | User selectable: 4, 8, 12 or 16 seconds |
| SpO₂ Data Update Period | 1 second |
| SpO₂ Alarm Condition Delay | 21 seconds (maximum time from physiological change to detection by device, with default 8 second averaging selected) |
| SpO₂ Alarm Signal Generation Delay | 1 second (time for device to generate alarm after alarm condition is detected) |
| SpO ₂ Sensitivity | User selectable: Normal, High |
| SpO ₂ Measurement | Functional SpO ₂ values are displayed and stored |
| Pulse Rate Accuracy (spe | ecified for range 25 to 240 bpm) |
| Adults/Pediatrics (RMS) | ±3% (during no motion conditions) ±5% (during motion conditions) |
| | |

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Specifications and Performance Characteristics

| | display with autogain control |
|---|---|
| SpCO™ | |
| SpCO Concentration Display Range | 0 to 40% |
| SpCO Accuracy (RMS)* | ±3% (during no motion conditions) |
| SpMet™ | |
| SpMet Saturation Range | 0 to 15.0% |
| SpMet Display Resolution | 0.1% up to 10% |
| SpMet Accuracy (RMS)* | ±1% (during no motion conditions) |
| about two-thirds of the mo above. Note: A functional tester o | e, SpCO, and SpMet measurements are statistically distributed, only easurements can be expected to fall within the accuracies specified cannot be used to assess SpO ₂ accuracy. See IEC 80601-2-61 |
| Annex FF. | |
| SpO₂ Measurement Waveleng Note: Information about w performing photodynamic | vavelength range can be useful to clinicians, for example, when |
| Masimo (SpO ₂ only) | Red: 660 nanometers |
| | Infrared: 905 nanometers |
| Nellcor (SpO₂ only) | Red: 660 nanometers Infrared: 900 nanometers |
| SpO ₂ Optical Power | |
| Masimo | Maximum optical output power = 15 mW (SpO ₂ only) Maximum optical output power for Rainbow sensor (SpO ₂ , SpCO, SpMet) = 25 mW |
| Nellcor | Maximum optical output power = 15 mW (SpO ₂ only) |
| NIBP | |
| Blood Pressure | Systolic Pressure Range: 30 to 255 mmHg |
| | Diastolic Pressure Range: 15 to 220 mmHg |
| • | Mean Arterial Pressure Range: 20 to 235 mmHg |
| | |
| | Units: mmHg in increments of 1 mmHg |
| | Units: mmHg in increments of 1 mmHg Blood Pressure Accuracy: ±5 mmHg |
| | • |
| Pulse Rate | Blood Pressure Accuracy: ±5 mmHg Blood Pressure Measurement Time: 20 seconds, typical (excluding |
| Pulse Rate Operation Features | Blood Pressure Accuracy: ±5 mmHg Blood Pressure Measurement Time: 20 seconds, typical (excluding cuff inflation time) Pulse Rate Range: 30 to 240 pulses per minute Pulse Rate Accuracy: ±2 pulses per minute or ±2%, whichever is |
| Operation Features | Blood Pressure Accuracy: ±5 mmHg Blood Pressure Measurement Time: 20 seconds, typical (excluding cuff inflation time) Pulse Rate Range: 30 to 240 pulses per minute Pulse Rate Accuracy: ±2 pulses per minute or ±2%, whichever is greater Initial Cuff Pressure: User selectable, 80 to 180 mmHg |

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Appendix A | Specifications and Performance Characteristics

| 2 | | | | | |
|------------------------------|---|--|--|--|--|
| CO₂ Range | 0 to 99 mmHg (0 to 13.2 kPa) Units: mmHg, %, or kPa | | | | |
| CO₂ Accuracy* | CO ₂ partial pressure at sea level: | | | | |
| (0–80 bpm)** | 0 to 38 mmHg (0 to 5.1 kPa) | ±2 mmHg (0.27 kPa) | | | |
| | 39 to 99 mmHg (5.2 to 13.2 kPa) | ±5% of reading + 0.08% for every 1 mmHg (0.13 kPa) above 38 mmHg (5.1 kPa) | | | |
| (>80 bpm) | 0 to 18 mmHg (0 to 2.4 kPa) | ±2 mmHg (0.27 kPa) | | | |
| | 19 to 99 mmHg (2.55 to 13.3 kPa) | ± 4 mmHg (0.54 kPa) or $\pm 12\%$ of reading, whichever is higher | | | |
| | *Determined by the methods in ISO 21647 clauses 51.101.1 and 51.101.2. **For RR > 60 bpm, to achieve specified CO₂ accuracy, the Microstream® FilterLine® H Set for infant must be used. | | | | |
| | The accuracy specifications listed above are maintained within an additional 4% in the presence of interfering gases defined in ISO 80601-2-55. | | | | |
| Respiration Rate Accuracy | 0 to 70 bpm: ±1 bpm 71 to 99 bpm: ±2 bpm | | | | |
| Respiration Rate Range | 0 to 99 breaths/minute | | | | |
| Flow Rate | 50 ml/min -7.5, +15 ml/min (flow measured by volume) | | | | |
| Rise Time | 190 msec | | | | |
| Response Time | 3.3 seconds with 200 cm FilterLine tubing 5.5 seconds with 400 cm FilterLine tubing (includes delay time and rise time) | | | | |
| Initialization Time | 30 seconds (typical), 10-180 seconds | | | | |
| Ambient Pressure | Automatically compen | sated internally | | | |
| Optional Display Waveform | CO₂ pressure | | | | |
| Scale Factors | Autoscale, 0-20 mmHg (0-4 Vol%),0-50 mmHg (0-7 Vol%), 0-100 mmHg (0-14 Vol%) | | | | |
| Waveform Sample Rate | 20/sec or one sample | every 50 msec | | | |
| CO₂ Calculation | Per 80601-2-55, Method for End-tidal CO₂ calculation: | | | | |
| | EtCO₂ is a maximum rather than average value. | | | | |
| | The accuracy of the Co with a CO ₂ square way | O₂ readings and respiration rates was tested e simulator. | | | |
| Measurement Drift | The periodic autozero function compensates for drifts between components, and changes in ambient temperature and barometric conditions. This automatic process eliminates variances that might otherwise cause measurement drift. Therefore, the CO ₂ function does not exhibit drift. | | | | |

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Specifications and Performance Characteristics

| Transducer Type | Strain-gauge resistive bridge |
|------------------------------|---|
| Transducer Sensitivity | 5μV/V/mmHg required |
| Defibrillation Protection | Minimum 360 J defibrillation protection required in the transducer |
| Excitation Voltage | 5 Vdc |
| Connector | Electro Shield CXS 3102A 14S-6S |
| Bandwidth | Digital filtered, DC to 30 Hz (< -3db) |
| Zero Drift | 1 mmHg/hr without transducer drift |
| Zero Adjustment | ±150 mmHg including transducer offset |
| Numeric Accuracy | ±1 mmHg or 2% of reading, whichever is greater, plus transducer error |
| Pressure Range | -30 to 300 mmHg, in six user selectable ranges |
| Invasive Pressure Display | Display: IP waveform and numerics Units: mmHg Labels: P1 or P2, ART, PA, CVP, ICP, LAP (user selectable) |
| TEMPERATURE | |
| Sensors | Measurement Specialties 4400 series esophageal/rectal and Foley catheter temperature probes, and 4499HD skin temperature probe |
| Displayed Range | 24.8° to 45.2°C (76.6° to 113.4°F) |
| Resolution | 0.1°C |
| Accuracy | ±0.2°C |
| Labels | Temp, T-esoph, T-naso, T-bladder, T-rectal, T-skin |
| Update Rate | Every 10 seconds, minimum |
| Mode of Operation | Direct mode |
| Adapter Cable | Only use Physio-Control part number 3303935 |
| Cable Length | 1.5 or 3 m (5 or 10 ft) |
| TREND | |
| Time Scale | Auto, 30 minutes, 1, 2, 4, or 8 hours |
| Duration | Up to 8 hours |
| ST | After initial 12-lead ECG analysis, automatically selects and trends ECG lead with the greatest ST displacement |
| Display | Choice of HR, PR (SpO ₂), PR (NIBP), SpO ₂ (%), SpCO(%), SpMet(%) CO ₂ (EtCO ₂ /FiCO ₂), RR (CO ₂), NIBP, IP1, IP2, TEMP, ST |
| NTERPRETIVE ALGORITHM | 12-Lead Interpretive Algorithm: University of Glasgow 12-Lead ECG Analysis Program, includes AMI and STEMI statements |
| ALARMS | |
| Quick Set | Activates alarms for all active vital signs |
| VF/VT Alarm | Activates continuous CPSS monitoring in Manual mode |
| Apnea Alarm | Occurs when 30 seconds has elapsed since last detected respiration |
| Heart Rate Alarm Limit | Upper, 100–250 bpm; lower, 30–150 bpm |

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Appendix A | Specifications and Performance Characteristics

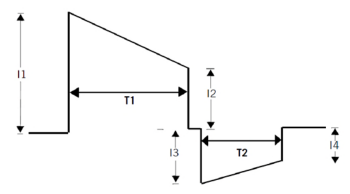
| PRINTER | | | | | |
|-------------|---|---|--|--|--|
| | ntinuous strip o | f the displayed patient info | ormation and reports | | |
| | Paper Size 100 mm (3.9 in) | | | | |
| | Print Speed 25 mm/sec or 12.5 m | | sec le base for 12-lead ECG reports | | |
| Delay | | 8 seconds | | | |
| Autoprint | : | Waveform events print a | automatically | | |
| Frequenc | cy Response | Diagnostic – 0.05 to 150 Hz or 0.05 to 40 Hz Monitor – 0.67 to 40 Hz or 1 to 30 Hz | | | |
| DEFIBRILLA | TOR | | | | |
| Charge Time | e (per IEC 60601 | -2-4) | | | |
| AC Operatio | n Only: | | | | |
| | Maximum Ti | me from Charge to Shock | Ready (Manual Mode): | | |
| | Voltage | | Charge Time | | |
| | 90-240 V | ac | 360 J within 10 seconds | | |
| | Maximum Ti | me from Initiation of Analy | rsis to Shock Ready (AED Mode): | | |
| | Voltage | | Charge Time | | |
| | 90-240 Vac | | 360 J within 30 seconds | | |
| | Maximum Ti | me from Power-on to Sho | ck Ready (Manual Mode): | | |
| | Voltage | | Charge Time | | |
| | 90-240 Vac | | 360 J within 25 seconds | | |
| | Maximum Time from Power-on to | | ck Ready (AED Mode): | | |
| | Voltage | | Charge Time | | |
| | 90-240 Vac | | 360 J within 40 seconds | | |
| DC Operatio | n Only: | | | | |
| | Maximum Ti | me from Charge to Shock | Ready (Manual Mode): | | |
| | Voltage | | Charge Time | | |
| | 11-17.6 Vdc | | 360 J within 10 seconds | | |
| | Maximum Time from Initiation of Analysis to Shock Ready (AED Mode): | | | | |
| | Voltage | | Charge Time | | |
| | 11-17.6 Vdc | | 360 J within 30 seconds | | |
| | Maximum Ti | me from Power-on to Sho | ck Ready (Manual Mode): | | |
| | Voltage | | Charge Time | | |
| | 11-17.6 V | /dc | 360 J within 25 seconds | | |
| | Maximum Ti | me from Power-on to Sho | ck Ready (AED Mode): | | |
| | Voltage | | Charge Time | | |
| | 11-17.6 V | /dc | 360 J within 40 seconds | | |
| | | | | | |

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Specifications and Performance Characteristics

| | Maximum 7 | ime from Charge to Sho | ock Ready (Manual Mode): | | | | |
|--|---|---|--|--|--|--|--|
| | Battery Status | | Charge Time | | | | |
| | Fully charged | | 200 J within 7 seconds, nominal | | | | |
| | | arged, followed by 15 | 360 J within 10 seconds | | | | |
| | Fully ch | arged | 360 J within 10 seconds nalysis to Shock Ready (AED Mode): Charge Time | | | | |
| | Maximum 1 | ime from Initiation of An | | | | | |
| | Battery | Status | | | | | |
| | Fully ch | arged | 200 J within 15 seconds, nominal | | | | |
| | | arged, followed by 15 gy shocks | 360 J within 30 seconds 360 J within 30 seconds | | | | |
| | Fully ch | arged | | | | | |
| | Maximum 7 | ime from Power-on to S | Shock Ready (Manual Mode): | | | | |
| | Battery | Status | Charge Time | | | | |
| | Fully charged, followed by 15 full-energy shocks | | 360 J within 25 seconds | | | | |
| | Maximum Time from Power-on to S Battery Status Fully charged, followed by 15 full-energy shocks | | Shock Ready (AED Mode): | | | | |
| | | | Charge Time | | | | |
| | | | 360 J within 40 seconds | | | | |
| anual Mode | | | | | | | |
| Energy Se | lect | 2, 3, 4, 5, 6, 7, 8, 9, 1 225, 250, 275, 300, 3 | 0, 15, 20, 30, 50, 70, 100, 125, 150, 175, 200, 25, and 360 joules | | | | |
| Synchronous cardioversion | | The maximum time delay between synchronization pulse and the delivery of energy, once the output has been activated, is not more than 60 msec. This time delay is measured from the peak of the QR to the peak of the defibrillator waveform. | | | | | |
| Sensing Leads Off if the retain 300 $\pm 15\%\Omega$ | | Leads Off if the resist | DMBO electrodes, the device indicates Paddles tive part of the patient impedance is greater if the magnitude of the patient impedance is $\%\Omega$. | | | | |
| Biphasic V | Vaveform | Biphasic Truncated Exponential | | | | | |
| | | The following specifications apply from 25 to 200Ω , unless otherwise specified: | | | | | |
| | | | Energy Accuracy: ± 1 joule or 10% of setting, whichever is greater, into $50\Omega_{;}$ ± 2 joules or 15% of setting, whichever is greater, into 25-175 $\Omega_{:}$ | | | | |
| | | Voltage Compensation: Active when disposable therapy electrodes are attached. Energy output within $\pm 5\%$ or ± 1 joule, whichever is greater, of 50Ω value, limited to the available energy which results in the delivery of 360 joules into 50Ω . | | | | | |

Waveform Shape and Measured Parameters

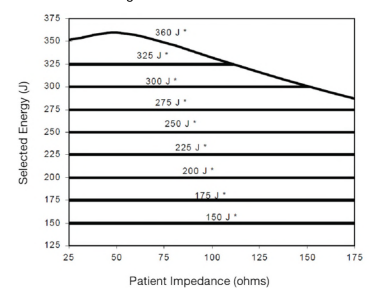


Biphasic Waveform

| Patient | Phase 1 Duration (ms) | | Phase 2 Du | ıration (ms) | Tilt (%) | |
|------------------------|-----------------------|------|------------|--------------|----------|------|
| Impedance (Ω) | Min | Max | Min | Max | Min | Max |
| 25 | 5.1 | 6.0 | 3.2 | 4.2 | 69.9 | 85.2 |
| 50 | 6.8 | 7.9 | 4.4 | 5.5 | 57.0 | 74.7 |
| 75 | 7.6 | 9.4 | 4.9 | 6.5 | 49.3 | 67.6 |
| 100 | 8.7 | 10.6 | 5.6 | 7.3 | 43.0 | 62.2 |
| 125 | 9.5 | 11.2 | 6.2 | 7.7 | 39.0 | 56.6 |
| 150 | 10.1 | 11.9 | 6.6 | 8.2 | 36.8 | 52.6 |
| 175 | 10.6 | 12.5 | 6.9 | 8.6 | 33.8 | 49.3 |

Rated Energy Output

Rated energy output is the nominal delivered energy based on the energy setting and patient impedance, as defined in the following chart.



^{*}Energy setting selected

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Specifications and Performance Characteristics

| Paddle Options | QUIK-COMBO pacing/defibrillation/ECG electrodes (standard) Standard paddles (optional) |
|---|--|
| Cable Length | 8 foot long (2.4 m) QUIK-COMBO cable (not including electrode assembly) |
| AED Mode | Shock Advisory System (SAS) is an ECG analysis system that advises the operator if the algorithm detects a shockable or nonshockable ECG rhythm. SAS acquires ECG via therapy electrodes only. |
| Biphasic Output Energy | Shock levels ranging from 150–360 joules with same or greater energy level for each successive shock |
| cprMAX™ Technology | In AED mode, cprMAX technology provides a method of maximizing the CPR time that a patient receives, with the overall goal of improving the rate of survival of patients treated with AEDs |
| Setup Options: | |
| Auto Analyze | Allows for auto analysis. Options are OFF, AFTER 1ST SHOCK |
| Initial CPR | Allows the user to be prompted for CPR for a period of time prior to other activity. Options are OFF , ANALYZE FIRST , CPR FIRST |
| Initial CPR Time | Time interval for Initial CPR. Options are 15, 30, 45, 60, 90, 120, and 180 seconds |
| Pre-Shock CPR | Allows the user to be prompted for CPR while the device is charging. Options are OFF , 15 , 30 seconds |
| Pulse Check | Allows the user to be prompted for a pulse check at various times. Options are ALWAYS, AFTER SECOND NSA, AFTER EVERY NSA, NEVER |
| Stacked Shocks | Allows for CPR after 3 consecutive shocks or after a single shock. Options are OFF , ON |
| CPR Time 1 or 2 | User selectable times for CPR. Options are 15, 30, 45, 60, 90, 120, 180 seconds and 30 minutes |
| PACER | |
| Pacing Mode | Demand or nondemand Rate and current defaults |
| Pacing Rate | 40 to 170 PPM |
| Rate Accuracy | ±1.5% over entire range |
| Output Waveform | Monophasic, truncated exponential current pulse (20 ±1 msec) |
| Output Current | 0 to 200 mA |
| | Pause: Pacing pulse frequency reduced by a factor of 4 when activated |
| Output Current Accuracy | \pm 10% or 5 mA (whichever is greater) over the specified load impedance range |
| Refractory Period | 180 to 280 msec (function of rate) |
| Physio-Control Therapy Electrode Post-Pacing Performance per IEC 60601-2-4 | After pacing: AC large signal impedance ≤ 4.2 Ω DC offset voltage ≤ 1053 mV After pacing followed by 360 J shock: DC offset voltage ≤ 1228 mV, 4 seconds after shock DC offset voltage ≤ 966 mV, 60 seconds after shock |

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Appendix A | Specifications and Performance Characteristics

| Operating Temperature | 0° to 45°C (32° to 113°F) -20°C (-4°F) for 1 hour after storage at room temperature 60°C (140°F) for 1 hour after storage at room temperature |
|------------------------------------|---|
| Storage Temperature | -20° to 65°C (-4° to 149°F) except therapy electrodes and batteries |
| Relative Humidity, Operating | 5 to 95%, non-condensing NIBP: 15 to 95%, non-condensing |
| Relative Humidity, Storage | 10 to 95%, non-condensing |
| Atmospheric Pressure, Operating | -382 to 4,572 m (-1,253 to 15,000 ft) NIBP: -152 to 3,048 m (-500 to 10,000 ft) |
| Water Resistance, Operating | IP44 (dust and splash resistance) per IEC 60529 and EN 1789 (without accessories except for 12-lead ECG cable, hard paddles, and battery pack) |
| Vibration | MIL-STD-810E Method 514.4 Propeller Aircraft - category 4 (figure 514.4-7 spectrum a) Helicopter - category 6 (3.75 Grms) Ground Mobile - category 8 (3.14 Grms) |
| | EN 1789: Sinusoidal Sweep, 1 octave/min, 10-150 Hz, ±0.15 mm/2 g |
| Shock (drop) | 5 drops on each side from 18 inches onto a steel surface EN 1789: 30-inch drop onto each of 6 surfaces |
| Shock (functional) | Meets IEC 60068-2-27 and MIL-STD-810E shock requirements 3 shocks per face at 40 g, 6 ms half-sine pulses |
| Bump | 1000 bumps at 15 g with pulse duration of 6 msec |
| Impact, Non-operating | IEC 60601-1 0.5 + 0.05 joule impact UL 60601-1 6.78 Nm impact with 2-inch diameter steel ball Meets IEC 62262 protection level IK04 |
| EMC | IEC 60601-1-2 Medical Equipment - General Requirements for Safety - Collateral Standard: Electromagnetic Compatibility - Requirements and Tests |
| Cleaning | Cleaning 20 times with the following: Quaternary ammonium, isopropyl alcohol, hydrogen peroxide |
| Chemical Resistance | 60 hour exposure to specified chemicals: Betadine (10% Povidone-Iodine solution) Coffee, Cola Dextrose (5% Glucose solution) Electrode Gel/Paste (98% water, 2% Carbopol 940) HCL (0.5% solution, pH=1) Isopropyl Alcohol NaCI solution (0.9% solution) Cosmetic discoloration of the paddle well shorting bar shall be |

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Specifications and Performance Characteristics

ESSENTIAL PERFORMANCE

The LIFEPAK 15 monitor/defibrillator includes the following essential performance features:

- Defibrillation, Synchronized Cardioversion, and AED Shock Advisory System
- ECG Monitoring, Heart Rate, and Alarms
- SpO₂ Monitoring, Pulse Rate, and Alarms
- EtCO₂ Monitoring and Alarms
- NIBP Monitoring and Alarms
- Invasive Pressure Monitoring and Alarms
- Temperature Monitoring and Alarms

Table 40 Battery Specifications

| CHARACTERISTIC | DESCRIPTION |
|---|--|
| Battery Type | Lithium-ion |
| Weight | < 0.60 kg (1.3 lb) |
| Charge Time (with fully depleted battery) | < 190 minutes (typical) |
| Battery indicators | Each battery has a fuel gauge that indicates its approximate charge. A fuel gauge that shows two or fewer LEDs after a charge cycle indicates that the battery should be replaced. |
| Charging Temperature Range | 5° to 45°C (41° to 113°F) |
| Operating Temperature Range | 0° to 45°C (32° to 113°F) |
| Short Term (<1 week) Storage Temperature Range | -20° to 60°C (-4° to 140°F) |
| Long Term (>1 week) Storage Temperature Range | 20° to 25°C (68° to 77°F) |
| Operating and Storage Humidity Range | 5 to 95% relative humidity, non-condensing |

Appendix A | Specifications and Performance Characteristics

Table 41 Alarm Limits

| Table 41 Alarn | n Limits | | | | | | | | |
|-----------------------------------|---------------|----------|----------|----------|-------------|--------|------------|--------|--------------|
| VITAL SIGN (VS) | PATIENT VS | WIDE I | _IMITS* | | ROW ITS* | LIMITS | RANGE | | AULT TS** |
| (٧3) | VALUE | LOW | HIGH | LOW | HIGH | LOW | HIGH | LOW | HIGH |
| Heart Rate | <60 | -20 | +35 | -10 | +25 | 30–150 | 100–250 | 50 | 150 |
| (HR) | 60–79 | -25 | +40 | -20 | +30 | | | | |
| Pulse Rate (PR) | 80–104 | -30 | +40 | -30 | +30 | | | | |
| (bpm) | ≥105 | -35 | +45 | -25 | +25 | | | | |
| SpO ₂ | ≥90 | -5 | +3 | -5 | +3 | 50 | 90–100 | 85 | 100 |
| (%) | <90 | -5 | +3 | -5 | +3 | | | | |
| | <90 | -20 | +35 | -10 | +25 | 30 | 245 | 50 | 200 |
| Systolic BP | 90–114 | -20 | +35 | -10 | +25 | | | | |
| (mmHg) | 115–140 | -25 | +35 | -10 | +20 | | | | |
| | >140 | -25 | +35 | -10 | +20 | | | | |
| D: . !! DD | <65 | -15 | +25 | -10 | +25 | 12 | 210 | 20 | 150 |
| Diastolic BP (mmHg) | 65-90 | -15 | +15 | -15 | +10 | | | | |
| | >90 | -15 | +15 | -15 | +10 | | | | |
| EtCO ₂ | >40/5.3 | -10/-1.3 | +15/+2.0 | -10/-1.3 | +15/+2.0 | 5/0.7 | 70/9.2 | 15/2.0 | 50/6.6 |
| (mmHg/%) | ≤40/5.3 | -10/-1.3 | +15/+2.0 | -10/-1.3 | +15/+2.0 | | | | |
| Inspired CO ₂ (mmHg/%) | _ | n/a | +5/+0.7 | n/a | +3/+0.4 | n/a | 0/0–10/1.3 | _ | 8/1.1 |
| Respiration | <15 | -8 | +8 | -4 | +4 | 5-15 | 10–60 | 5 | 30 |
| Rate (RR) | ≥15 | -15 | +15 | -8 | +8 | | | | |
| | <15 | -6 | +12 | -4 | +6 | 10 | 100 | 10 | 40 |
| Systolic PA (mmHg) | 15–35 | -8 | +16 | -6 | +8 | | | | |
| (11111119) | >35 | -12 | +16 | -8 | +10 | | | | |
| D: . !! DA | <5 | -4 | +12 | -4 | +8 | 0 | 50 | 0 | 18 |
| Diastolic PA (mmHg) | 5–13 | -4 | +16 | -6 | +6 | | | | |
| | >13 | -6 | +16 | -6 | +6 | | | | |
| CVP (mmHg) | ≥9 | -10 | +10 | -5 | +5 | 0 | 25 | 0 | 15 |
| ICP, LAP | <15 | -6 | +6 | -4 | +4 | 0 | 40 | 0 | 18 |
| (mmHg) | ≥15 | -6 | +8 | -4 | +6 | | | | |
| Temperature (°C) | ≥31 | -3 | +3 | -1 | +1 | 31 | 41 | 35 | 39 |

^{*}Numbers are \pm from patient's VS value when the alarms are set.

^{**}Default limits are established when alarms are set up to be ON.

Specifications and Performance Characteristics

| Table 42 Alarm Performa | ance Characteristics | | | | |
|-------------------------|---|--|--|--|--|
| CHARACTERISTIC | DESCRIPTION | | | | |
| Heart Rate Alarm Time | For a 1 mV, 206 bpm tachycardia, the average detection time was 4.6 seconds. | | | | |
| | For a test signal half as large, the average was 4.1 seconds. In this case the device sensitivity was increased to 5mV/cm. | | | | |
| | For a test signal twice as large, the average was 3.1 seconds. | | | | |
| | For a 2 mV, 195 bpm tachycardia, the average detection time was 2.5 seconds. | | | | |
| | For a test signal half as large, the average was 2.2 seconds. In this case the device sensitivity was increased to 5mV/cm. | | | | |
| | For a test signal twice as large, the average was 1.5 seconds. | | | | |
| Audible Alarms | This is a standalone device. All alarm tones are internal to the LIFEPAK 15 monitor/defibrillator. The alarm tone volumes range from 45 to 85 dB. | | | | |
| | Alarm violations are manifested by tones, voice prompts, and visual indications. | | | | |
| | Alarm manifestation occurs within 1 second after a displayed parameter violates its alarm limit. User selectable alarm volume adjustment is provided. This adjustment does not allow alarm volume to attain/reach a zero level. | | | | |
| | SAS tones reinforce SAS messages provided on the product display. | | | | |
| | The following identifies the tone assignments for each type of alarm: | | | | |
| | The priority 1 tone is used to alert the user to the possibility of death. This tone is a 440 Hz and 880 Hz alternating tone with a 50% duty cycle and a 4 Hz alternation frequency. This tone has a volume of 70 ±5 dB (A) as measured at a distance of 1 meter from the display. The volume of the priority 1 alarm is not adjustable. | | | | |
| | The priority 2 tone (the Quick Set alarm tone) is used to alert the user that a possible life-threatening condition exists. This tone is a continuous 698 Hz tone. This tone has a volume that is lower than the priority 1 tone. | | | | |
| | The priority 3 tone is used to alert the user that an abnormal condition exists. Three beeps at 1046 Hz for 100 ms duration each with a 150 ms silence between the first and second and the second and third, followed by a 200 ms silence. This tone has a volume that is lower than the priority 2 tone. | | | | |
| | Priority 3 tones come in single and repeating types: for a single tone, the 3-beep sequence sounds only once. For a repeating tone, the 3- beep sequence sounds every 20 seconds. | | | | |
| | The priority 4 tone is a momentary tone between 500 and 1500 Hz. This tone has a volume that is lower than the priority 3 tone. Specific characteristics are: | | | | |
| | QRS and Volume Setting Tone—100 msec duration at 1397 Hz— | | | | |

4 msec duration at 1319 Hz.

Appendix A | Specifications and Performance Characteristics

| | PECADIDEION |
|----------------------|---|
| CHARACTERISTIC | DESCRIPTION |
| | The alert tone shall consist of one set of two tones to precede voice prompts and to draw attention to the display. Specific characteristics consist of: |
| | 1000 Hz square wave, 100 ms duration |
| | Silence, 100 msec duration |
| | 1000 Hz square wave, 100 ms duration |
| | Silence, 140 msec duration (when preceding a voice prompt) |
| | Voice prompt, when used |
| Visual Alarms | Alarms are indicated visually by: |
| | The violated parameter flashes in inverse video with a message in the message area of the display. |
| | These visual indications remain on the display until the alarm is corrected. Visual indication of alarms continue even when the tones have been silenced. |
| Priorities for Alarm | Alarm conditions have the following priorities. |
| Conditions | Priority 1: |
| | VF or VT detected based on ECG signal |
| | Priority 2: |
| | Heart rate high or low limit exceeded |
| | SpO₂ high or low limit exceeded |
| | NIBP high or low limit exceeded |
| | Invasive pressure high or low limit exceeded |
| | EtCO₂ high or low limit exceeded |
| | FiCO₂ high limit is exceeded |
| | Respiration rate high or low limit exceeded |
| | Apnea detected |
| | Temperature high or low limit exceeded |
| | Priority 3: |
| | ECG leads off detected |
| | Low battery (5-15% capacity) detected |
| | Very low battery (<5% capacity) detected Note: For this condition, the priority 3 tone is followed by the REPLACE BATTERY prompts. |
| | Service condition that could prevent normal operation detected. |
| | |

Specifications and Performance Characteristics

Table 43 Setup Options Factory Default Settings

| General Repeate (a country Specific) (a cole Summary Confect (a cole Summary Col Summary Cole Summary Cole Summary Cole Summary Cole Summary Col Summary Cole Summary C | MENU | Options Factory Defaumental MENU/ITEM | FACTORY DEFAULT SETTINGS | |
|---|-------------|---------------------------------------|--------------------------|--------------------|
| Trend Summary Off Site Number 000 Device ID "LP15" + last 4 digits of serial number, for example, LP151234 Auto Log On Line Filter 60 Hz Timeout Speed 30 seconds Manual mode Sync After Shock Off Pads Default 200 (joules) Energy Protocol Inactive Internal Default 10 (joules) Voice Prompts On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200-300-360 Auto Analyze Off Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds Metronome On Metronome On Off Mult - No Airway 30:2 Of | General | Language | (Country Specific) | |
| Site Number 000 Device ID "LP15" + last 4 digits of serial number, for example, LP151234 Auto Log On Line Filter 60 Hz Timeout Speed 30 seconds Manual mode Sync After Shock Off Pads Default 200 (joules) Energy Protocol Inactive Internal Default 10 (joules) Yoice Prompts On Manual Access Manual / Direct Set Passcode 000 Manual Access Manual / Direct Set Passcode 000 Auto Analyze Off Motion Detection On Auto Analyze Off Motion Detection On Pulse Check Never CPR Time 1 120 seconds Initial CPR Time 1 120 seconds Initial CPR Time 2 120 seconds Preshock CPR Off Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 10:2 | | Code Summary | Long | |
| Device ID "LP15" + last 4 digits of serial number, for example, LP151234 Auto Log On Line Filter 60 Hz Timeout Speed 30 seconds Manual mode Sync After Shock Off Pads Default 200 (joules) Energy Protocol Inactive Internal Default 10 (joules) Voice Prompts On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200-300-360 Auto Analyze Off Motion Detection On Pulse Check Never CPR Never CPR Time 1 120 seconds Initial CPR Off Metronome Adult - No Ainway 10:1 Youth - No Ainway 10:1 Youth - No Ainway 10:1 Youth - No Ainway | | Trend Summary | Off | |
| Line Filter | | Site Number | 000 | |
| Line Filter 60 Hz Timeout Speed 30 seconds Manual mode Sync After Shock Off Pads Default 200 (joules) Energy Protocol Inactive Internal Default 10 (joules) Voice Prompts On Shock Tone On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200-300-360 Auto Analyze Off Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off Metronome Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 10:1 Youth - No Airway 10:1 Youth - No Airway 10:1 Youth - Airway 10:1 Youth - No Airway 10:1 Youth - No Airway 10:1 | | Device ID | | mber, for example, |
| Manual mode Manual mode Metronome Sync After Shock Off Off Manual mode Mode Mode Mode Mode Mode Mode Mode | | Auto Log | On | |
| Manual mode Angle A | | Line Filter | 60 Hz | |
| Pads Default 200 (joules) Energy Protocol Inactive Internal Default 10 (joules) Voice Prompts On Shock Tone On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200–300–360 Auto Analyze Off Motion Detection On Pulse Check Never CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Channel 2 </td <td></td> <td>Timeout Speed</td> <td>30 seconds</td> <td></td> | | Timeout Speed | 30 seconds | |
| Energy Protocol Inactive Internal Default 10 (joules) Voice Prompts On Shock Tone On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200–300–360 Auto Analyze Off Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Adult - No Airway 30:2 Adult - No Airway 10:1 Youth - Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Monitoring Channels Default Set None | Manual mode | Sync After Shock | Off | |
| Internal Default 10 (joules) Voice Prompts On | | Pads Default | 200 (joules) | |
| Voice Prompts On Shock Tone On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200-300-360 Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds Energy Protocol CPR Time 2 120 seconds CPR Time 2 120 seconds Initial CPR Time 120 seconds Initial CPR Time 120 seconds Preshock CPR Off Metronome On Multiple Properties On Packed Adult - No Airway 30:2 Adult - No Airway 15:2 Youth - No Airway 15:2 Youth - No Airway On H Packed On PM Mode Demand | | Energy Protocol | Inactive | |
| Shock Tone On Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200-300-360 Auto Analyze Off Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds Energy Protocol CPR Time 2 120 seconds CPR Time 2 120 seconds Initial CPR Time 120 seconds Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Metronome On Metronome On Multi- No Airway 30:2 Adult - No Airway 15:2 Youth - No Airway 15:2 Youth - No Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Default Set Set 1 Monitoring Channel 2 None | | Internal Default | 10 (joules) | |
| Manual Access Manual / Direct Set Passcode 0000 AED mode Energy Protocol 200–300–360 Auto Analyze Off Motion Detection On Pulse Check Never CPR Time 1 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off Metronome On < | | Voice Prompts | On | |
| Set Passcode 0000 AED mode Energy Protocol 200–300–360 Auto Analyze Off Motion Detection On Pulse Check Never CPR Time 1 120 seconds Initial CPR Time 2 120 seconds Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Metronome On Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Channel 2 None Channel 3 None | | Shock Tone | On | |
| AED mode Energy Protocol 200–300–360 Auto Analyze Off Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Channel 2 None Channel 3 None | | Manual Access | Manual / Direct | |
| Auto Analyze Off Motion Detection On Pulse Check Never CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Set Passcode | 0000 | |
| Motion Detection On Pulse Check Never CPR CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off Metronome On Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | AED mode | Energy Protocol | 200–300–360 | |
| Pulse Check Never CPR CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off CPR Metronome On Metronome On Off Adult - No Airway 30:2 Off Adult - Airway 10:1 Off Youth - No Airway 15:2 Off Youth - Airway 10:1 Off Pacing Rate 60 PPM Current 0 mA Off Mode Demand Off Internal Pacer Detection Off Monitoring Channels Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Auto Analyze | Off | |
| CPR CPR Time 1 120 seconds CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off CPR Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 2 None Channel 3 None | | Motion Detection | On | |
| CPR Time 2 120 seconds Initial CPR Off Initial CPR Time 120 seconds Preshock CPR Off CPR Metronome Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Pulse Check | Never | |
| Initial CPR | | CPR | CPR Time 1 | 120 seconds |
| Initial CPR Time 120 seconds | | | CPR Time 2 | 120 seconds |
| Preshock CPR Off CPR Metronome Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | | Initial CPR | Off |
| CPR Metronome Metronome On Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | | Initial CPR Time | 120 seconds |
| Metronome Adult - No Airway 30:2 Adult - Airway 10:1 Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | | Preshock CPR | Off |
| Adult - No Airway 10:1 | _ | Metronome | On | |
| Youth - No Airway 15:2 Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | Metronome | Adult - No Airway | 30:2 | |
| Youth - Airway 10:1 Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Adult - Airway | 10:1 | |
| Pacing Rate 60 PPM Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Youth - No Airway | 15:2 | |
| Current 0 mA Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Youth - Airway | 10:1 | |
| Mode Demand Internal Pacer Detection Off Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | Pacing | Rate | 60 PPM | |
| Monitoring Detection Off Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Current | 0 mA | |
| Monitoring Channels Default Set Set 1 Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Mode | Demand | |
| Set 1 Channel 1 ECG Lead II Channel 2 None Channel 3 None | | Internal Pacer | Detection Off | |
| Channel 2 None Channel 3 None | Monitoring | Channels | Default Set | Set 1 |
| Channel 3 None | | Set 1 | Channel 1 | ECG Lead II |
| | | | Channel 2 | None |
| Continuous Data ECG Channel 1 | | | Channel 3 | None |
| | | Continuous Data | ECG Channel 1 | |

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Appendix A | Specifications and Performance Characteristics

| MENU | MENU/ITEM | FACTORY DEFAULT SETTINGS | |
|---------|-----------------------|-----------------------------|---------------|
| | SpO ₂ Tone | Off | |
| | CO ₂ | Units | mmHg |
| | | BTPS | Off |
| | NIBP | Initial Pressure | 160 mmHg |
| | | Interval | Off |
| | Temperature | Units | Celsius |
| | Trends | On | |
| 12-Lead | Auto Transmit | Off | |
| | Auto Print | On | |
| | Print Speed | 25 mm/sec | |
| | Interpretation | On | |
| | Format | 3-Channel Standard | |
| Events | Events Page 1 | Event 2 | Oxygen |
| | | Event 3 | IV Access |
| | | Event 4 | Nitroglycerin |
| | | Event 5 | Morphine |
| | | Event 6 | Cancel Last |
| | | Event 7 | Intubation |
| | | Event 8 | CPR |
| | | Event 9 | Epinephrine |
| | | Event 10 | Atropine |
| | | Event 11 | Lidocaine |
| | Events Page 2 | Event 12 | ASA |
| | | Event 13 | Heparin |
| | | Event 14 | Thrombolytic |
| | | Event 15 | Glucose |
| | | Event 16 | Naloxone |
| | | Event 17 | Transport |
| | | Event 18 | Adenosine |
| | | Event 19 | Vasopressin |
| | | Event 20 | Amiodarone |
| | | Event 21 | Dopamine |
| | | Event 22 | Bicarb |
| Alarms | Volume | 5 | |
| | Alarms | Off | |
| | VF/VT Alarm | Off | |

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Specifications and Performance Characteristics

| MENU | MENU/ITEM | FACTORY DEFAULT SETTINGS | |
|--------------|--------------------------------|------------------------------------|-----|
| Printer | Auto Print | Defibrillation | On |
| | | Pacing | Off |
| | | Check Patient | Off |
| | | SAS | Off |
| | | Patient Alarms | Off |
| | | Events | Off |
| | | Initial Rhythm | Off |
| | ECG Mode | Monitor | |
| | Monitor Mode | 1–30 Hz | |
| | Diagnostic Mode | .05–40 Hz | |
| | Alarm Waveforms | On | |
| | Event Waveforms | On | |
| | Vitals Waveforms | Off | |
| Transmission | Sites | Site 1 / Output Port / Direct Conn | ect |
| | Default Site | None | |
| | Default Report | 12-Lead | |
| | Wireless | On | |
| | Search Filter | On | |
| Clock | Date/Time | Current date/time PST | |
| | Clock Mode | Real Time | |
| | DST | Off | |
| | Time Zone | None | |
| Self Test | Transmit Results | Off | |
| Service | Maintenance Prompt Interval | Off | |

Appendix B

Screen Messages

This appendix describes the screen messages that the LIFEPAK 15 monitor/defibrillator may display during normal operation.

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Summary of Screen Messages

Table 44 Summary of Screen Messages

| MESSAGE | DESCRIPTION |
|----------------------------|--|
| 12-LEAD ECG UNAVAILABLE | A 12-lead was requested but the necessary ECG data is not available. |
| ABNORMAL ENERGY DELIVERY | A discharge occurred when the paddles were shorted together, when hard paddles did not have adequate contact with the patient or were discharged in the air, or patient impedance was out of range. Message may also appear in certain types of internal faults. |
| ACCESS DENIED | Three consecutive incorrect passcode attempts were made to enter Manual mode. |
| ACQUIRING 12-LEAD | Monitor is acquiring data for 12-lead ECG report. |
| ACQUIRING SNAPSHOT | A snapshot report of current vital signs has been requested. |
| ADVISORY MODE-MONITORING | The device is monitoring the patient ECG for a shockable rhythm. |
| ADVISORY: SPCO > 10% | SpCO advisory alert activated. SpCO value is greater than 10%. |
| ADVISORY: SPMET > 3% | SpMet advisory alert activated. SpMet value is greater than 3%. |
| ALARM APNEA | No valid breath has been detected for 30 seconds. |
| ALARMS SILENCED | Alarms are silenced. An alert tone with status message ALARMS SILENCED occurs periodically as a reminder. |
| ANALYZING 12-LEAD | The data for 12-lead ECG report is being analyzed. |
| ANALYZING NOW-STAND CLEAR | The AED is analyzing the patient ECG rhythm. |
| ATTEMPTING TO TRANSMIT | The device is processing a transmission request. |
| AUTO NIBP CANCELLED | The automatic initiation of NIBP measurements has been cancelled. |
| BATTERY X LOW | The specified battery has a low energy condition. |
| BLUETOOTH DEVICE NOT FOUND | Bluetooth device has not been detected. |
| BLUETOOTH UNAVAILABLE | Unable to locate or connect to target device. |
| CANNOT CHARGE | CHARGE is pressed and the synchronize source is missing for synchronized cardioversion, the therapy cable is not connected, or QUIK-COMBO electrodes are not attached to the therapy cable. |
| CANNOT CHARGE BATTERIES | Both batteries are installed, and the device is unable to charge either battery. |
| CANNOT CHARGE BATTERY 1 | The device is unable to charge the battery in battery well 1. |
| CANNOT CHARGE BATTERY 2 | The device is unable to charge the battery in battery well 2. |
| CHARGING TO XXX J | Appears when CHARGE is pressed on the front panel or standard paddles. |
| CHECK FOR PULSE | AED prompt after each standard 3-shock sequence or NO SHOCK ADVISED message. |
| | A potentially shockable rhythm has been detected when the |

Summary of Screen Messages

| MESSAGE | DESCRIPTION |
|---|---|
| CHECK PATIENT. IF NO PULSE, PUSH ANALYZE | A potentially shockable rhythm has been detected when using Advisory Monitoring. |
| CHECK PRINTER | The printer door is open, there is no paper in the printer, or another printer malfunction exists. |
| CO2 AUTOZERO | EtCO ₂ monitor is automatically performing a zero-point calibration. |
| CO2 FILTERLINE BLOCKAGE | EtCO ₂ FilterLine tubing is kinked or clogged; the message appears after 30 seconds of unsuccessful purging. |
| CO2 FILTERLINE OFF | EtCO ₂ FilterLine tubing is disconnected or is not securely connected to the device. |
| CO2 FILTERLINE PURGING | EtCO ₂ FilterLine tubing is kinked or clogged with liquid. |
| CO2 INITIALIZING | EtCO₂ monitor is performing a self-check. |
| CONNECT CABLE | Therapy cable is not connected when you press CHARGE , PACER , or ANALYZE . |
| CONNECT CHEST LEADS | A 12-lead ECG analysis was requested and precordial leads are not connected to the patient. |
| CONNECT ECG LEADS | ECG electrodes or leads are disconnected. |
| CONNECT ELECTRODES | Therapy electrodes are disconnected. |
| CONNECTED TO | The device is connected via <i>Bluetooth</i> technology to another <i>Bluetooth</i> -enabled device. The name of the connected device follows this message. |
| CONNECTING TO | The device is establishing communication with another Bluetooth-enabled device. The name of the target device follows this message. |
| CPR: ADULT-AIRWAY X:Y | An option for CPR metronome. The patient is an adult for whom an advanced airway has been established. The specified C:V ratio will be used. |
| CPR: ADULT-NO AIRWAY X:Y | An option for CPR metronome. The patient is an adult for whom an advanced airway has not been established. The specified C:V ratio will be used. |
| CPR: YOUTH-AIRWAY X:Y | An option for CPR metronome. The patient is a youth (younger than the age of puberty) for whom an advanced airway has been established. The specified C:V ratio will be used. |
| CPR: YOUTH-NO AIRWAY X:Y | An option for CPR metronome. The patient is a youth (younger than the age of puberty) for whom an advanced airway has not been established. The specified C:V ratio will be used. |
| CURRENT FAULT | The comparison between delivered and selected pacing current is out of tolerance. |
| DEMAND | Pacemaker is in Demand mode. |
| DEMO MODE | The device is in Demo mode and simulated patient data is displayed. |
| DISARMING | The energy charge is being removed internally. |
| ECG CABLE OFF | The device is printing and the ECG cable is removed. |
| ECG LEADS OFF | Multiple ECG electrodes are disconnected. |
| ENDING DEVICE SEARCH | The request for finding a <i>Bluetooth</i> device was stopped. |
| ENERGY DELIVERED | Energy transfer is complete. |

Appendix B | Screen Messages

| MESSAGE | DESCRIPTION |
|---|--|
| ENERGY FAULT | The comparison between stored and selected energy is out of tolerance. |
| ENTER MANUAL MODE? | One of the Manual mode access buttons was pressed and the confirmation screen is set up to appear. |
| EXCESSIVE NOISE - 12-LEAD CANCELLED | Noise is detected for longer than 30 seconds that is too great to record a 12-lead ECG report. |
| IF NO PULSE, PUSH ANALYZE | Follows a CPR interval, if a PULSE CHECK setup option other than NEVER is selected. |
| IF NO PULSE, START CPR | Follows delivery of a shock or NO SHOCK ADVISED prompt, if a PULSE CHECK setup option other than NEVER is selected. |
| IF YOU WITNESSED THE ARREST, PUSH ANALYZE | Initial CPR message that follows START CPR prompt, to remind user to deliver a shock immediately if the user witnessed the arrest. |
| LA LEADS OFF | ECG electrode "LA" is disconnected. |
| LAST CONNECTED TO | When <i>Bluetooth</i> connectivity is installed and this device previously connected to a target device, the name of the target device appears after this message. |
| LL LEADS OFF | ECG electrode "LL" is disconnected. |
| LOST BLUETOOTH CONNECTION | Communication with Bluetooth device has been interrupted. |
| LOST DIRECT CONNECTION | Communication via direct connection has been interrupted. |
| MAINTENANCE DUE | Reminder message that appears at the interval that is set in Service mode. Message continues to appear until reset or turned off. |
| MANUAL MODE DISABLED | Access to Manual mode from AED mode has been restricted. |
| MOTION DETECTED!/STOP MOTION! | Motion was detected during ECG analysis. |
| NIBP AIR LEAK | NIBP cuff applied too loosely or there is a leak in cuff/monitor pneumatic system. |
| NIBP CHECK CUFF | NIBP cuff is not connected to patient or device. |
| NIBP FAILED | NIBP monitor cannot establish zero-pressure reference. |
| NIBP FLOW ERROR | NIBP pneumatic system is not maintaining stable cuff pressure. |
| NIBP INITIALIZING | NIBP requested while NIBP module is still initializing. |
| NIBP MOTION | Patient extremity moved too much for the NIBP monitor to accurately complete the measurement. |
| NIBP OVERPRESSURE | NIBP cuff pressure exceeded 290 mmHg. |
| NIBP TIME OUT | NIBP monitor did not complete a measurement in 120 seconds. |
| NIBP WEAK PULSE | The monitor did not detect any pulses. |
| NO SHOCK ADVISED | The defibrillator did not detect a shockable rhythm. |
| NO SITES DEFINED | Device is attempting to transmit using <i>Bluetooth</i> connection, but no associated destinations have been defined. |
| NOISY DATA! PRESS 12-LEAD TO ACCEPT | Monitor detects excessive signal interference while acquiring data. Press 12-LEAD to override the message and acquire 12-lead ECG with noise. |
| NON-DEMAND | Pacemaker is in Nondemand (asynchronous) mode. |
| | |

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Summary of Screen Messages

| MESSAGE | DESCRIPTION | |
|---|---|--|
| OBTAINING DEVICE NAMES | Device is obtaining names of available <i>Bluetooth</i> -enabled devices. | |
| PACER FAULT | Internal error detected during pacing. | |
| PACING IN PROGRESS | The requested action is not available because the device is currently performing pacing. | |
| PACING STOPPED | Pacing has stopped—for example, due to disconnection of therapy electrodes. | |
| PASSCODE INCORRECT - TRY AGAIN | Incorrect passcode entered. | |
| PAUSED | The pacing PAUSE button is pressed and held. Current pulses are applied at reduced frequency while the MA and PPM settings are maintained. | |
| PUSH ANALYZE | Press ANALYZE to begin ECG analysis. | |
| PUSH AND HOLD SHOCK BUTTON! | The defibrillator is in Sync mode, fully charged, and ready to provide therapy. | |
| PUSH AND HOLD PADDLE BUTTONS TO SHOCK! | The defibrillator is in Sync mode, fully charged, and ready to provide therapy with hard paddles connected. | |
| PUSH SHOCK BUTTON! | The defibrillator is fully charged and ready to provide therapy. | |
| PX NOT ZEROED | Transducer is connected or reconnected and is not zeroed. | |
| PX TRANSDUCER NOT DETECTED | IP transducer is disconnected from the monitor/defibrillator. | |
| PX ZERO FAILED | The device was unable to zero the pressure transducer. | |
| PX ZEROED | Transducer successfully zeroed. | |
| PX ZEROING | Monitor is establishing a zero reference. | |
| RA LEADS OFF | ECG electrode "RA" is disconnected. | |
| REPLACE BATTERY X | Power loss for the battery in well X is imminent. | |
| SEARCHING FOR DEVICES | Device is attempting to identify available <i>Bluetooth</i> devices. | |
| SELECT BIPHASIC ENERGY / XXX J | ENERGY SELECT was pressed on front panel or on standard paddles. | |
| SELF TEST FAILED | Device detected internal error; remove device from service. | |
| SELF TEST FAILED. TRANSMITTING | Device detected internal error and is transmitting test results. Remove device from service after transmission is complete. | |
| SELF TEST IN PROGRESS | Device is performing a self test after turning on. | |
| SELF TEST PASSED | Device passed internal test and is available for use. | |
| SELF TEST PASSED. TRANSMITTING | Device passed internal test and is transmitting test results. | |
| SHOCK ADVISED! | The defibrillator has analyzed the patient ECG rhythm and detected a shockable ECG rhythm. | |
| SPCO: POOR QUALITY SIGNAL | Device is not receiving sufficient input from sensor. | |
| SPMET: POOR QUALITY SIGNAL | Device is not receiving sufficient input from sensor. | |
| SPO2: CHECK SENSOR | The SpO₂ sensor connection to device or application to patient needs checked. | |
| SPO2: LOW PERFUSION | Patient has a weak pulse. | |
| SPO2: NO SENSOR DETECTED | A sensor is disconnected from the monitor. | |
| SPO2: POOR QUALITY SIGNAL | Device is not receiving sufficient input from sensor. | |

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Appendix B | Screen Messages

| MESSAGE | DESCRIPTION |
|---|---|
| SPO2: SEARCHING FOR PULSE | A sensor is connected to the patient and is searching for a |
| SFO2. SEARCHING FOR PULSE | pulse. |
| SPO2: SENSOR DOES NOT SUPPORT SPCO OR SPMET | The sensor in use only measures SpO ₂ . |
| SPO2: UNKNOWN SENSOR | A sensor that is not Physio-Control approved is connected to the device. |
| STAND CLEAR/PUSH SHOCK BUTTON | Prompts you to stand clear and push 🗲 (shock). |
| START CPR | Prompts you to begin providing CPR to the patient. |
| SWITCHING PRIMARY TO LEAD II | Pacing is turned on while PADDLES is the primary lead. |
| SWITCHING PRIMARY TO PADDLES | Device was in Lead II when ANALYZE was pressed. PADDLES becomes the primary lead. |
| SYNC MODE | Device is currently in Sync mode. |
| TEMP: ACCURACY OUTSIDE LIMITS | Temperature accuracy check has failed. |
| TEMP: CHECK SENSOR | Device is not receiving sufficient input from sensor. |
| TO CANCEL, PUSH SPEED DIAL | The defibrillator is charging or charged and the device may be disarmed by pressing the Speed Dial. |
| TRANSMISSION CANCELLED | Data transmission has been cancelled. |
| TRANSMISSION COMPLETED | Data transmission completed successfully. |
| TRANSMISSION FAILED | Data transmission was not successful. |
| TRANSMITTING TO <site></site> | Connection is established to <site> and transmission of requested report is occurring.</site> |
| UNABLE TO CONNECT | Unable to establish connection with Bluetooth device. |
| UNABLE TO TRANSMIT | Unable to send data. |
| UNKNOWN DEVICE | Bluetooth connection failed or timed out before obtaining target device name. |
| USE ECG LEADS | Sync mode attempted, but ECG electrodes are not attached to patient, PADDLES lead is displayed, and standard paddles are connected to defibrillator. |
| USER TEST FAILED | Unsuccessful User Test. |
| USER TEST IN PROGRESS | USER TEST selected on the OPTIONS menu and test is in process. |
| USER TEST PASSED | Successful User Test completed. |
| VX LEADS OFF | ECG electrode such as "V1" is disconnected. |
| X DEVICES FOUND | Shows number of <i>Bluetooth</i> -enabled devices found. |
| XX LEADS OFF | ECG electrode such as "RA" is disconnected. |
| XX% TRANSMITTED | Specified percent of the transmission is completed. |

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Appendix C

Shock Advisory System

This appendix describes the basic function of the Shock Advisory System™ (SAS) algorithm.

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Overview of the Shock Advisory System

The Shock Advisory System (SAS™) is an ECG analysis system built into the biphasic LIFEPAK 15 monitor/defibrillator that advises the operator as to whether it detects a shockable or nonshockable rhythm. This system makes it possible for individuals who are not trained to interpret ECG rhythms to provide potentially lifesaving therapy to victims of ventricular fibrillation or pulseless ventricular tachycardia.

The Shock Advisory System contains the following features:

- Electrode Contact Determination
- Automated Interpretation of the ECG
- Operator Control of Shock Therapy
- Continuous Patient Surveillance System (CPSS)
- Motion Detection

The Shock Advisory System is active when the LIFEPAK 15 monitor/defibrillator is used as an automated external defibrillator (AED). CPSS may be activated during monitoring.

Upon the user pressing the ₹ (shock) button, the LIFEPAK 15 monitor/defibrillator delivers the shock therapy to the patient.

Electrode Contact Determination

The Shock Advisory System measures the patient's transthoracic impedance through the therapy electrodes. If the baseline impedance is higher than a maximum limit, it determines that the electrodes do not have sufficient contact with the patient or are not properly connected to the AED. When this occurs, ECG analysis and shock delivery are inhibited. The AED advises the operator to connect electrodes when there is insufficient electrode contact.

Automated Interpretation of the ECG

The Shock Advisory System recommends a shock if it detects the following:

- Ventricular fibrillation—with a peak-to-peak amplitude of at least 0.08 mV.
- Ventricular tachycardia—defined as having a heart rate of at least 120 beats per minute, QRS width of at least 0.16 seconds, and no apparent P waves.

Pacemaker pulses may prevent advisement of an appropriate shock, regardless of the patient's underlying rhythm. The Shock Advisory System recommends no shock for all other ECG rhythms including asystole, pulseless electrical activity, idioventricular rhythms, bradycardia, supraventricular tachycardias, atrial fibrillation and flutter, heart block, premature ventricular complexes, and normal sinus rhythms. These rhythms are specifically mentioned in the AHA recommendations. The SAS does not continue analyzing the ECG after a **SHOCK ADVISED** decision is reached.

Shock Advisory System Performance

ECG analysis by the Shock Advisory System (SAS) in the LIFEPAK 15 monitor/defibrillator was tested by playing ECG waveforms from the Physio-Control database through the electrode connector. For each test ECG, the decision **SHOCK** or **NO SHOCK** of the SAS was recorded and compared to the rhythm classification and treatment recommendation by clinical experts. A report of test results is available on request.

SAS Test Set

The SAS Test Set consists of 989 ECG samples recorded during pre-hospital use of the LIFEPAK 5 defibrillator. The ECG was recorded using cassette tape recorders connected to the LIFEPAK 5 defibrillator. Selected ECG segments were sampled and the ECG rhythm was classified by clinical experts. The SAS Test Set contains the following ECG samples:

- 168 each coarse ventricular fibrillation (VF) (≥200 µV peak-to-peak amplitude)
- 29 each fine ventricular fibrillation (<200 and ≥80 μV peak-to-peak amplitude)
- 65 each shockable ventricular tachycardia (VT) (HR >120 bpm, QRS duration ≥160 ms, no apparent P waves, patient reported to be pulseless by the paramedics)
- 43 each asystole (<80 μV peak-to-peak amplitude)
- 144 each normal sinus rhythm (NSR) (sinus rhythm, heart rate 60-100 bpm)
- 531 each other organized rhythm (includes all rhythms except those in other listed categories)
- 2 each transitional (transition occurs within the sample from nonshockable to shockable or vice versa)
- 5 each shockable rhythms with pacemaker artifact (the pacemaker artifact is spread over time by the filtering in the LIFEPAK 5 defibrillator)
- 2 each nonshockable rhythms with pacemaker artifact (the pacemaker artifact is spread over time by the filtering in the LIFEPAK 5 defibrillator)

Table 45 LIFEPAK 15 Monitor/Defibrillator Overall SAS Performnce

| >90% |
|------|
| >95% |
| >90% |
| <5% |
| |

Table 46 LIFEPAK 15 Monitor/Defibrillator Performance by Rhythm Category

| RHYTHM CLASS | ECG TEST ¹ SAMPLE SIZE | PERFORMANCE GOAL | OBSERVED PERFORMANCE |
|---------------------------------|--|-----------------------------------|---|
| Shockable: Coarse VF | 168 | >90% sensitivity | LIFEPAK 15 monitor/defibrillator meets the IEC 60601-2-4 requirements and AHA ² recommendations. |
| Shockable: VT | 65 | >75% sensitivity | LIFEPAK 15 monitor/defibrillator meets the IEC 60601-2-4 requirements and AHA recommendations. |
| Nonshockable: NSR | 144 | >99% specificity for NSR (AHA) | LIFEPAK 15 monitor/defibrillator meets the AHA recommendations. |
| Nonshockable: asystole | 43 | >95% specificity | LIFEPAK 15 monitor/defibrillator meets the IEC 60601-2-4 requirements and AHA recommendations. |
| Nonshockable: all other rhythms | 531 | >95% specificity | LIFEPAK 15 monitor/defibrillator meets the IEC 60601-2-4 requirements and AHA recommendations. |
| Intermediate: fine VF | 29 | Report only | >75% sensitivity |

¹ Each sample is run 10 times asynchronously.

Operator Control of Shock Therapy

The Shock Advisory System causes the AED to charge automatically when it detects the presence of a shockable rhythm. When a shock is advised, the operator presses the **SHOCK** button to deliver the energy to the patient.

Continuous Patient Surveillance System

The Continuous Patient Surveillance System (CPSS) automatically monitors the patient's ECG rhythm for a potentially shockable rhythm while the electrodes are attached and the AED is turned on. CPSS is not active during ECG analysis or when the AED is in a CPR cycle.

Motion Detection

The Shock Advisory System detects patient motion independent of ECG analysis. A motion detector is designed into the LIFEPAK 15 monitor/defibrillator. **MOTION DETECTION** can be set up to be **ON** or **OFF**. For more information, see the *LIFEPAK 15 Monitor/Defibrillator Setup Options* provided with your device.

A number of activities can create motion, including CPR, rescuer movement, patient movement, and some internal pacemakers. If variations in the transthoracic impedance signal exceed a maximum limit, the Shock Advisory System determines that patient motion of some kind is

² Automatic External Defibrillators for Public Access Defibrillation: Recommendations for Specifying and Reporting Arrhythmia Analysis Algorithm Performance, Incorporating New Waveforms, and Enhancing Safety. American Heart Association (AHA) Task Force on Automatic External Defibrillation, Subcommittee on AED Safety and Efficacy. Circulation, 1997: Vol. 95: 1677-1682.

VF = ventricular fibrillation VT = ventricular tachycardia

NSR = normal sinus rhythm

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Overview of the Shock Advisory System

present. If motion is detected, the ECG analysis is inhibited. The operator is advised by a displayed message, a voice prompt, and an audible alert. After 10 seconds, if motion is still present, the motion alert stops and the analysis always proceeds to completion. This limits the delay in therapy in situations where it may not be possible to stop the motion. However, the rescuer should remove the source of motion whenever possible to minimize the chance of artifact in the ECG.

There are two reasons why ECG analysis is inhibited when the motion alert occurs, and why the rescuer should remove the source of the motion whenever possible:

- 1. Such motion may cause artifact in the ECG signal. This artifact can cause a nonshockable ECG rhythm to look like a shockable rhythm. For example, chest compressions during asystole can look like shockable ventricular tachycardia. Artifact can also cause a shockable ECG rhythm to look like a nonshockable rhythm. For example, chest compressions during ventricular fibrillation can look like an organized, and therefore nonshockable, rhythm.
- 2. The motion may be caused by a rescuer's interventions. To reduce the risk of inadvertently shocking a rescuer, the motion alert prompts the rescuer to move away from the patient. This will stop the motion and ECG analysis will proceed.

Appendix D

SpO2 Clinical Validation Summaries

This appendix describes clinical validation data for SpO₂, SpCO, and SpMet monitoring.

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Masimo Clinical Validation Data

Masimo Corporation conducted clinical studies and tests to assess the accuracy of the SpO₂, SpCO, and SpMet measurement functions. The following paragraphs provide a summary of that data.

Test Methods for Accuracy

SpO₂, SpCO and SpMet accuracy was determined by testing on healthy adult volunteers in the range of 60-100% SpO₂, 0-40% SpCO, and 0-15% SpMet against a laboratory CO-oximeter. SpO₂ and SpMet accuracy was determined on 16 neonatal intensive care patients ranging in age from 7-135 days old and weighing between 0.5-4.25 kg. Seventy-nine (79) data samples were collected over a range of 70-100% SaO₂ and 0.5-2.5% MetHb with a resultant accuracy of 2.9% SpO₂ and 0.9% SpMet.

The Masimo sensors have been validated for no motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies in the range of 70-100% SpO₂ against a laboratory CO-oximeter and ECG monitor. This variation equals plus or minus one standard deviation. Plus or minus one standard deviation encompasses 68% of the population weight.

The Masimo sensors have been validated for motion accuracy in human blood studies on healthy adult male and female volunteers with light to dark skin pigmentation in induced hypoxia studies while performing rubbing and tapping motions at 2 to 4 Hz at an amplitude of 1 to 2 cm, and a non-repetitive motion between 1 to 5 Hz at an amplitude of 2 to 3 cm in induced hypoxia studies in the range of 70-100% SpO₂ against a laboratory CO-oximeter and ECG monitor. This variation equals plus or minus one standard deviation which encompasses 68% of the population.

The Masimo sensors have been validated for pulse rate accuracy for the range of 25-240 bpm in bench top testing against a Biotek Index 2 simulator. This variation equals plus or minus one standard deviation which encompasses 68% of the population.

Nellcor Clinical Validation Data

Nellcor Clinical Validation Data

Nellcor conducted clinical studies and tests to assess the accuracy of the SpO₂ measurement function of the Nellcor SpO₂ sensors. The following paragraphs provide a summary of that data.

Test Methods for Accuracy

 SpO_2 accuracy specifications for Nellcor sensors are based on controlled hypoxia studies with healthy nonsmoking adult volunteers over the specified saturation SpO_2 range. Pulse oximeter SpO_2 readings were compared to SaO_2 values of drawn blood samples measured by hemoximetry.

Subjects used to validate SpO2 measurement accuracies were healthy and recruited from the local population. The study group comprised both men and women; subjects spanned a range of skin pigmentations and ranged in age from 18-50 years old. When sensors are used on neonatal subjects as recommended, the specified accuracy is decreased by $\pm 1\%$, as compared to adult usage, to account for the theoretical effect on oximeter measurements of fetal hemoglobin in neonatal blood.

Appendix E

Electromagnetic Compatibility Guidance

This appendix provides guidance and manufacturer's declaration of electromagnetic compatibility.

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Electromagnetic Compatibility Guidance

Electromagnetic Emissions

Table 47 Guidance and Manufacturer's Declaration - Electromagnetic Emissions

The LIFEPAK 15 monitor/defibrillator is intended for use in the electromagnetic environment specified below. The customer or the user of the LIFEPAK 15 monitor/defibrillator should assure that it is used in such an environment.

| Emissions Test | Compliance | Electromagnetic Environment - Guidance | |
|--|----------------|---|--|
| RF emissions CISPR 11 | Group 1 | The LIFEPAK 15 monitor/defibrillator uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. | |
| RF emissions CISPR 11 | Class B | The LIFEPAK 15 monitor/defibrillator is suitable for use in all establishments, including domestic establishments and | |
| Harmonic emissions IEC 61000-3-2 | Not applicable | those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes. | |
| Voltage fluctuations/ flicker emissions IEC 61000-3-3 | Not applicable | | |

Electromagnetic Compatibility Guidance

Electromagnetic Immunity

Table 48 Guidance and Manufacturer's Declaration - Electromagnetic Immunity

The LIFEPAK 15 monitor/defibrillator is intended for use in the electromagnetic environment specified below. The customer or the user of the LIFEPAK 15 monitor/defibrillator should assure that it is used in such an environment.

| Immunity Test | IEC 60601 Test Level | Compliance Level | Electromagnetic Environment - Guidance |
|---|--|---|--|
| Electrostatic discharge (ESD) IEC 61000-4-2 | ±6 kV contact ±8 kV air | ±6 kV contact ±8 kV air | Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%. |
| Electrical fast transient/burst IEC 61000-4-4 | ±2 kV for power supply lines ±1 kV for input/output lines | ±2 kV for power supply lines ±1 kV for input/output lines | Mains power quality should be that of a typical commercial or hospital environment. |
| Surge IEC 61000-4-5 | ±1 kV line(s) to line(s) ±2 kV line(s) to earth | ±1 kV line(s) to line(s) ±2 kV line(s) to earth | Mains power quality should be that of a typical commercial or hospital environment. |
| Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11 | $<5\%$ $U_{\rm T}$ (>95% dip in $U_{\rm T}$) for 0.5 cycle 40% $U_{\rm T}$ (60% dip in $U_{\rm T}$) for 5 cycles 70% $U_{\rm T}$ (30% dip in $U_{\rm T}$) for 25 cycles $<5\%$ $U_{\rm T}$ (>95% dip in $U_{\rm T}$) for 5 sec | $<5\% \ U_{T}$ $(>95\% \ dip \ in \ U_{T})$ for 0.5 cycle $40\% \ U_{T}$ $(60\% \ dip \ in \ U_{T})$ for 5 cycles $70\% \ U_{T}$ $(30\% \ dip \ in \ U_{T})$ for 25 cycles $<5\% \ U_{T}$ $(>95\% \ dip \ in \ U_{T})$ for 5 sec | Mains power quality should be that of a typical commercial or hospital environment. If the user of the LIFEPAK 15 monitor/defibrillator requires continued operation during power mains interruptions, it is recommended that the LIFEPAK 15 monitor/defibrillator be powered from an uninterruptible power supply or a battery. |
| Power frequency (50/60 Hz) magnetic field IEC 61000-4-8 | 3 A/m | 3 A/m | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment. |

Table 49 Guidance and Manufacturer's Declaration - Electromagnetic Immunity

The LIFEPAK 15 monitor/defibrillator is intended for use in the electromagnetic environment specified below. The customer or the user of the LIFEPAK 15 monitor/defibrillator should assure that it is used in such an environment.

| Immunity Test | IEC 60601 Test Level | Compliance Level | Electromagnetic Environment - Guidance |
|-------------------------------|--|---------------------|--|
| | | | Portable and mobile RF communications equipment should be used no closer to any part of the LIFEPAK 15 monitor/defibrillator, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. |
| | | | Recommended separation distance |
| Conducted RF IEC 61000-4-6 | 3 Vrms 150 kHz to 80 MHz outside ISM bands ¹ | 3 Vrms | $d = 1.2\sqrt{P}$ |
| | 10 Vrms 150 kHz to 80 MHz in ISM bands ¹ | 10 Vrms | $d=1.2\sqrt{P}$ |
| Radiated RF | 10 V/m 80 MHz to 2.5 GHz | 10 V/m | $d = 1.2\sqrt{P}$ 80 MHz to 800 MHz |
| IEC 61000-4-3 | | | $d = 2.3\sqrt{P}$ 800 MHz to 2.5 GHz |
| | | | Where <i>P</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in meters (m). ² Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, ³ should be less than the compliance level in each frequency range. ⁴ Interference may occur in the vicinity of equipment marked with the following symbol: |
| | | | equipment marked with the following symbol: |
| | | | ((2)) |

Note: At 80 MHz and 800 MHz, the higher frequency range applies.

Note: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

¹ The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

² The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz are intended to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in these frequency ranges.

³ Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the LIFEPAK 15 monitor/defibrillator is used exceeds the applicable RF compliance level above, the LIFEPAK 15 monitor/defibrillator should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the LIFEPAK 15 monitor/defibrillator.

⁴ Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Electromagnetic Compatibility Guidance

Separation Distances

Table 50 Recommended Separation Distances Between Portable and Mobile RF Communications Equipment and the LIFEPAK 15 Monitor/Defibrillator

The LIFEPAK 15 monitor/defibrillator is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the LIFEPAK 15 monitor/defibrillator can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the LIFEPAK 15 monitor/defibrillator as recommended below, according to the maximum output power of the communications equipment.

| Rated maximum output power of transmitter | Separation distance according to frequency of transmitter m | | | |
|---|---|--|-------------------------------------|--------------------------------------|
| | 150 kHz to 80 MHz outside ISM bands $d = 1.2\sqrt{P}$ | 150 kHz to 80 MHz in ISM bands $d = 1.2\sqrt{P}$ | 80 MHz to 800 MHz $d = 1.2\sqrt{P}$ | 800 MHz to 2.5 GHz $d = 2.3\sqrt{P}$ |
| 0.01 | 0.12 | 0.12 | 0.12 | 0.23 |
| 0.1 | 0.38 | 0.38 | 0.38 | 0.73 |
| 1 | 1.2 | 1.2 | 1.2 | 2.3 |
| 10 | 3.8 | 3.8 | 3.8 | 7.3 |
| 100 | 12 | 12 | 12 | 23 |

For transmitters rated at a maximum output power not listed above, the recommended separation distance *d* in meters (m) can be determined using the equation applicable to the frequency of the transmitter, where *P* is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

Note: The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

Note: An additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

Note: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Appendix F

Symbols

This appendix provides information about the symbols that are used in these operating instructions, or on the LIFEPAK 15 monitor/defibrillator, its accessories, packaging, or training tools.

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Symbols

The symbols in the following table may be found in these operating instructions or on the LIFEPAK 15 monitor/defibrillator, its accessories, packaging, or training tools.

Table 51 Symbols

| SYMBOL | DESCRIPTION |
|------------------------|--|
| Device or User Interfa | се |
| <u>i</u> | Attention, consult accompanying documents |
| | Follow instructions for use |
| | General warning |
| | Alarm on |
| * | Alarm off |
| | VF/VT alarm on |
| × | VF/VT alarm is on, but is silenced or suspended |
| | Battery in well, fully charged. For a description of all battery indicators, see Battery Status Indicators (on page 38). |
| 4 | Heart rate/pulse rate indicator |
| * | Bluetooth wireless technology |
| (x) | Shock count (x) on screen |
| Ø | Shock button on front panel or hard paddles |
| ₹ | Auxiliary power indicator |
| \$ | Battery charging indicator |
| ₽ | Service indicator |
| > | Greater than |
| < | Less than |
| J | Joules |

Symbols

| SYMBOL | DESCRIPTION |
|--------------------|---|
| ₩ | Display mode button |
| | Home Screen button |
| 4 | CO ₂ input |
| CO2 | CO₂ exhaust |
| \Leftrightarrow | Input/output |
| - * | Defibrillation-proof type CF patient connection |
| ┤ | Defibrillation protected, type BF patient connection |
| X | Do not dispose of this product in the unsorted municipal waste stream. Dispose of this product according to local regulations. See www.physio-control.com/recycling for instructions on disposing of this product. |
| 50 | Symbol for China RoHS indicating the Environmentally Friendly Use Period (EFUP) denoting the number of years before any substance is likely to leak out into the environment. |
| CE | Mark of conformity to applicable European Directives |
| ⊕ ∪s | Canadian Standards Association certification for Canada and the United States |
| c us | Intertek certification for Canada and the United States |
| IP44 | Enclosure ingress protection code per IEC 60529 |
| M _{YYYY} | Date of manufacture. Date may appear before, after, or below the figure. |
| EC REP | Authorized EC representative |
| MIN or PN | Manufacturer's identification number (part number) |
| SN | Serial number |
| REF | Reorder number |
| Rx Only or Rx Only | By prescription only |
| !USA | For USA audiences only |
| CAT | Catalog number |
| | Manufacturer |

Appendix F | Symbols

| SYMBOL | DESCRIPTION |
|----------------------------|---|
| C _{N13571} | Indicates that a product complies with applicable Australian ACMA standards |
| + | Positive terminal |
| | Negative terminal |
| | Fuse |
| | Battery |
| $\overline{\hspace{1cm}}$ | Power input |
| | Static-sensitive device. Static discharge may cause damage. |
| Reports | |
| \mathcal{V}^{Γ} | Biphasic defibrillation shock |
| 1 | Pace arrow, noninvasive pacing |
| ↔ | Pace arrow, internal pacing detection |
| | QRS sense marker |
| lacksquare | Event marker |
| Accessories | |
| CE | Mark of conformity to applicable European Directives |
| 71 . | Recognized component mark for the United States |
| c Fl °us | Recognized component mark for Canada and the United States |
| F© | Complies with (USA) Federal Communications Commission regulations |
| ((\(\frac{1}{4}\)) | Device includes RF transmitter |
| ҡ | Type BF patient connection |
| LOT YYWW | Lot number (batch code). YY (year) and WW (week) of manufacture. |
| IP44 | Enclosure ingress protection code per IEC 60529 |
| A or | Warning, high voltage |

Symbols

| SYMBOL | DESCRIPTION |
|---------------------------------------|---|
| (A) | CAUTION - FIRE HAZARD Do not disassemble, heat above 100°C (212°F), or incinerate battery |
| (X) | CAUTION - FIRE HAZARD Do not crush, puncture, or disassemble battery |
| Ω | Use By date shown: yyyy-mm-dd or yyyy-mm |
| | Indoor use only |
| LATEX | Item is latex free |
| Pb | Lead free |
| | Dispose of properly |
| 50°C - 122°F 0°C - 32°F | Store in a cool, dry location (0° to 50°C, 32° to 122°F) |
| 2 | Single use only |
| 2 = 2 | 2 electrodes in 1 package |
| 10 x 2 = 10 (2) | 10 packages in 1 shelf-pak |
| 5 x 10 (2) = 50 (2) | 5 shelf-paks in 1 case |
| | Shave patient skin |
| | Clean patient skin |
| A A A A A A A A A A A A A A A A A A A | Treatment |
| | Tear here |
| | Press electrode firmly onto patient |
| | Connect QUIK-COMBO cable |
| | Slowly peel back protective liner on electrode |

Appendix F | Symbols

| SYMBOL | DESCRIPTION |
|--|---|
| LIFERIC 1900, 1000 LIFERIC CEP Place LIFERIC REVESSOR defaultable | Do not use this pediatric QUIK-COMBO electrode on LIFEPAK 500, LIFEPAK 1000, LIFEPAK CR® Plus, or LIFEPAK EXPRESS® defibrillators |
| | For use on adults |
| | Not for use on adults |
| | For use on children up to 15 kg (33 lb) |
| | Not for use on children under 15 kg (33 lb) |
| | Remove label from battery |
| | Charge battery |
| | Insert battery in LIFEPAK 15 monitor/defibrillator |
| (+/< | Rechargeable battery |
| 2== | AC-DC power adapter |
| | DC-DC power adapter |
| 15 | For use with the LIFEPAK 15 monitor/defibrillator |
| \rightarrow | Power input |
| \longrightarrow | Power output |
| | DC voltage |
| ~ | AC voltage |
| Shipping carton | |
| <u> </u> | This end up |

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Symbols

| SYMBOL | DESCRIPTION |
|-----------------------------|---|
| | Fragile/breakable Handle with care |
| Ť | Protect from water |
| -20°C (140°F) | Recommended storage temperature -20° to 60°C (-4° to 140°F) |
| ₁₀ ⁹⁵ | Relative humidity range 10 to 95% |
| ॐ or ॐ | Recycle this item |

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LIFEPAK 15 Monitor/Defibrillator Operator's Checklist



This is a recommended checklist to use to inspect and test this monitor/defibrillator. Daily inspection and test is recommended. This form may be reproduced.

| Unit Serial No: | Location: | | | | | _ | |
|---|---|-------------|--------|--------|--------|------|------|
| Inchmedian | Recommended | Date | | | | | |
| Instruction | Corrective Action | Initials | | | | | |
| | | | √ each | box af | ter co | mple | ting |
| Inspect physical condition for: Foreign substances | Clean the device. | | | | | | |
| Damage or cracks | Contact a qualified service ter | chnician. | | | | | |
| 2. Inspect power source for: Broken, loose, or worn battery pins | Contact a qualified service tea | chnician. | | | | | |
| Damaged or leaking battery | Recycle or discard battery. | | | | | | |
| Spare battery available | Obtain fully charged spare ba | ittery. | | | | | |
| Damage to power adapter and cables | Contact a qualified service te | chnician. | | | | | |
| 3. Inspect ECG cable and cable port for | r: | | • | | | | |
| Cracking, damage, broken, or bent parts or pins | Replace ECG cable. If port is damaged, contact queservice technician. | ualified | | | | | |
| 4. Check ECG electrodes and therapy e | | | | , , | • | ı | 1 |
| Use By date | Replace if date passed. | | | | | | |
| Spare electrodes available | Obtain spare electrodes. | | | | | | |
| Damaged, opened package | Discard and replace electrodes. | | | | | | |
| 5. With batteries installed, disconnect f ON and observe for: | rom power adapter (if using), | , press | | | | | |
| Momentary illumination of self-test messages and LEDs, and speaker beep | If absent, contact a qualified stechnician. | service | | | | | |
| Two fully charged batteries | Replace low battery or charge battery using power adapter. | e installed | | | | | |
| Service indicator (🗡) | If illuminated, contact a qualif service technician. | ied | | | | | |

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| Instructions | Recommended Corrective Action | Date Initials | | | |
|---|---|-------------------------------|--|--|---|
| 6. With batteries installed, reconnect po | | heck for: | | | |
| Power adapter LED strip is illuminated Auxiliary power LED on device is | | | | | |
| illuminated Battery charging LED on device is illuminated or flashing | If absent, check batteries. If p persists, contact a qualified se technician. | | | | |
| 7. Perform QUIK-COMBO® therapy cabe (If this cable is not used with the defibrence) Disconnect and examine cable for cracking, damage, broken, or bent parts or pins. | | oy cable. | | | |
| Connect therapy cable to defibrillator and the Test Load. | If CONNECT ELECTRODES, PADDLES LEADS OFF, CONNECT CABLE, or ABNORMAL ENERGY DELIVERY message appears, replace therapy cable and repeat check. If problem continues, remove the defibrillator from use and contact a qualified service technician. | | | | |
| Select LEAD, then PADDLES. Select 200 JOULES and press CHARGE. Press (shock) button. | | | | | |
| Confirm ENERGY DELIVERED message appears. | If message does not appear, r therapy cable and repeat chec | | | | _ |
| Remove Test Load from cable and verify PADDLES LEADS OFF appears.** | If absent, contact a qualified stechnician. | service | | | |
| 8. Perform standard (hard) paddles che (If hard paddles are not used with the d Disconnect and examine cable for cracking, damage, broken, or bent parts or pins. Connect paddles to defibrillator. | | | | | |
| Examine for paddle surface pitting | Replace paddles, or clean page | ddles. | | | - |
| and presence of dried or wet gel. | | | | | - |
| Press LEAD. Select PADDLES. On paddles, turn ENERGY SELECT | If selected energy does not ch | nange or | | | - |
| dial to 10 JOULES.*** With paddles in paddle wells, press CHARGE button on paddle. | charging does not occur, obta paddles and repeat check. If p continues, remove the defibril from use and contact a qualifi service technician. | ain spare oroblem lator | | | |

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| Inotorotiono | Recommended Corrective Action | Date | | | | | |
|---|--|---|--|--|---|--|--|
| Instructions | | Initials | | | | | |
| | | | | | • | | |
| Press only one (shock) button and release. Confirm that energy was not discharged. | If energy discharges with one press, obtain spare paddles a repeat check. | | | | | | |
| Press the other (shock) button and release. Confirm that energy was not discharged. | | | | | | | |
| Press both (shock) buttons and confirm ABNORMAL ENERGY DELIVERY message appears. | If message does not appear, obtain spare paddles and repeat check. If problem continues, remove the defibrillator from use and contact a qualified service technician. | | | | | | |
| Remove paddles from wells, and confirm artifact on screen. | repeat check. If problem cont | If task fails, obtain spare paddles and repeat check. If problem continues, | | | | | |
| Place paddle surfaces together, and confirm flat line on screen. | remove the defibrillator from use and contact a qualified service technician. | | | | | | |
| Return paddles securely to paddle wells. | | | | | | | |
| 9. Perform User Test if 3:00 am auto test results not available: | | | | | | | |
| Press OPTIONS.Select USER TEST in menu.Confirm test results printed. | If User Test fails, remove the defibrillator from use and conqualified service technician. | tact a | | | | | |
| 10. Check ECG printer for: | | | | | | | |
| Adequate paper supply | Add new paper, if necessary. | | | | | | |
| Ability to print | If not working, contact a quali service technician. | fied | | | | | |
| 11. If using wireless data transmission, test transmission method: | | | | | | | |
| Establish a Bluetooth connection. | If not working, contact a quali service technician. | fied | | | | | |
| Send a test transmission. | | | | | | | |
| 12. Turn off defibrillator. (Press and hold ON for up to 2 seconds | .) | | | | | | |
| 13. Confirm that the device is stowed, n | nounted, or positioned secur | ely. | | | | | |

^{*} The defibrillator delivers up to 360 joules of electrical energy. Unless discharged properly, this electrical energy may cause serious personal injury or death. Do not attempt to perform this test unless you are qualified by training and experience.

^{**} Failure to remove the Test Load may result in delay of therapy during patient use.

^{***} Discharging > 10 joules in the paddle wells may damage the defibrillator.

LIFEPAK® 15 MONITOR/DEFIBRILLATOR

Operating Instructions

For further information, please contact your local Physio-Control representative or visit www.physio-control.com



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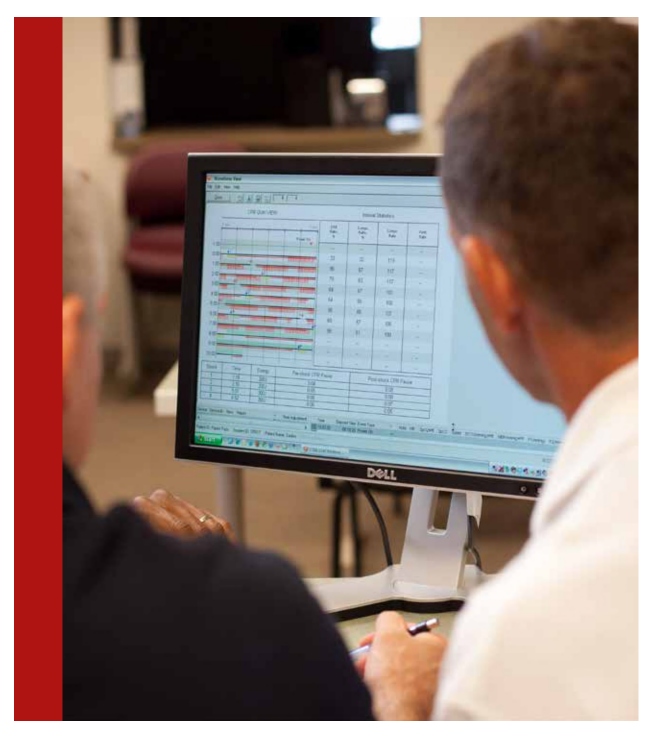
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EXHIBIT 2



CODE-STAT[™] 10

DATA REVIEW SOFTWARE



BASIC ANNOTATION HANDBOOK

Special thanks to Dana Yost, from Redmond Medic One, for writing and guidance support.

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Introduction

Welcome to CODE-STAT™ 10 data review software. This Basic Annotation Handbook is designed as a supplement to the CODE-STAT 10 user's guide. It provides a step-by-step guide for reviewing and annotating cases in CODE-STAT software, and utilizing the data for continuous quality improvement (CQI).

As you develop a CODE-STAT annotation system within your organization, pay particular attention to the number of annotators trained in relation to the number of cases available to annotate. Just like other technical duties, CODE-STAT annotation requires practice and frequency of use in order to stay proficient.

For detailed instructions regarding the use of the software, please refer to the CODE-STAT user's guide found by clicking ? or by going to **Help** \rightarrow **View Help**.

Before You Begin

When using the LIFEPAK® 12 and 15 monitor/defibrillators, defibrillation electrodes (pads) must be applied to the patient and the device must be in PADDLES lead in Channel 1 in order to record the impedance signal that is required to annotate cardiac arrest cases. Only defibrillation electrodes (pads) must be applied to patient in order to record the impedance channel for LIFEPAK 20/20e monitor/defibrillator and LIFEPAK 15 monitor/defibrillator with (1) icon on the label located on the back of the device in the battery well (see below).



In order to utilize the automated ventilation detection algorithm, your device must be capable of recording continuous capnography in CODE-STAT data review software. The LIFEPAK 12 monitor/defibrillator does not have this capability. Refer to LIFEPAK 15 monitor/defibrillator set up information on next page to determine if your device is capable.

Monitor/Defibrillator Set Up

Not all LIFEPAK 15 monitor/defibrillators have the ability to record the capnography waveform in CODE-STAT data review software. To determine if your 15 has this capability:

- 1. Go into the setup mode.
- 2. Select **Monitoring**. Check to see which of the setup options you have below:

If you see: Continuous ECG

Your device does not have the ability to record continuous capnography waveform data in CODE-STAT data review software and the automated ventilation detector will not be activated. You will still see the capnography waveform on the LIFEPAK screen if you have capnography; it just won't be recorded in the CODE-STAT data review software. Make sure the option is set to **ON**.

If you see: Continuous Data

Your device does have the ability to record continuous capnography waveform data in CODE-STAT data review software and the automated ventilation detector will be activated. You will still see the capnography waveform on the LIFEPAK screen if you have capnography and it will be recorded in the CODE-STAT data review software. Make sure the option is set to **All Channels**.

LIFEPAK 20/20e

In order to see the continuous capnography waveform in CODE-STAT software, you will need to:

- Enter the setup mode of monitor/defibrillator
- Select Monitoring
- Select Continuous ECG
- Select ON

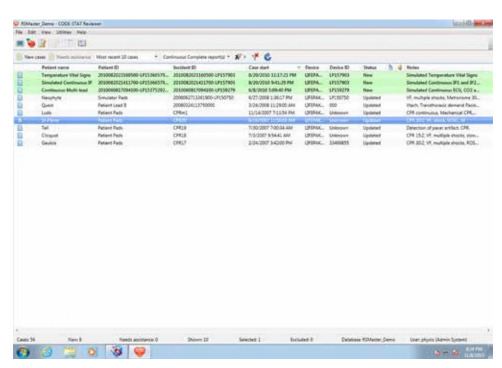
Getting Familiar with CODE-STAT Data Review Software

For the basic tour of CODE-STAT data review software, we will use illustrations from the demo database included when the product was installed. Launch CODE-STAT software by selecting (or double clicking) **CODE-STAT Reviewer**.

When the Database Login screen appears, enter the following demo database credentials:

> User ID: physio Password: control

Once logged in, you will see the case list view, which is used for navigating through cases.



The following case list fields are displayed:

📄 Hide/show case details.

Patient Name The patient's last name (if entered).

Patient ID The patient ID, either automatically assigned by the device or

entered in the field. The default ID is a date and time stamp

along with the device serial number.

Incident ID Same as above.

Case Start The case start time and date.

Device The type of the device used in collecting the data.

Device ID The ID assigned to that device.

Status The status of the case. This can be helpful during the

annotation and editing process.

This icon is used to indicate a file attachment.

This icon is used to indicate cases with audio files.

Notes Additional notes that can be entered regarding the case.

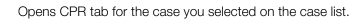
You can sort the columns by clicking the column header.

Top Level Buttons—Case List





Opens the highlighted case.





Opens the data entry window for the highlighted case in the case list view.

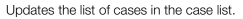
Displays help.



A flashing green button indicates new cases. Click to display the new cases.



Indicates that there are cases that need assistance due to import issues.





Hides opened case details for all cases on list.



Runs DT EXPRESS.



Allows for time-based filtering of the case list.



Allows for filtering by report type.



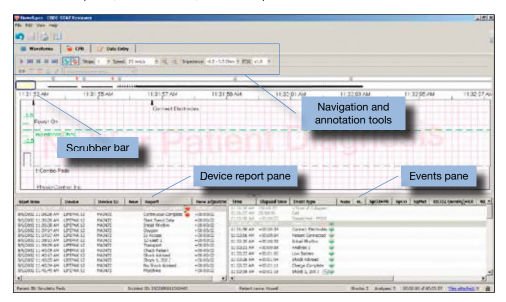
Opens Text search filter.



Clears currently selected filter criteria to display all cases on the list.

Opening a Case

Open a case by double clicking it in the case list view, or highlighting it and then clicking . This will bring you to the case view where you can see the continuous waveform reports, 12-lead ECG, or other reports and associated events.



You can navigate through a case in several ways. The buttons at the top of the page allow you to either play the entire case, or move by page or by segment (this is covered in more detail in the annotation section). Alternatively, you can use the horizontal scroll (scrubber) bar to move through the case. Simply click in the yellow box and move your mouse to the right.

Clicking on any event in the events pane will navigate you to the location of the event in the continuous ECG report.

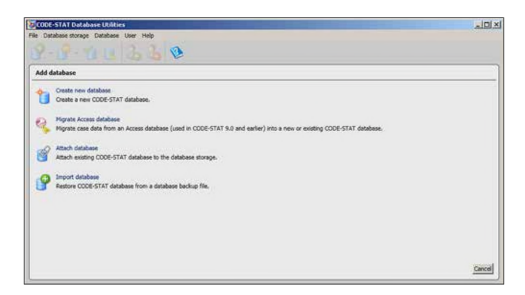
Set Up a New Database

New Database

Before you get started working with real cases, it is important to set up a new database. This allows you to keep your cases outside of the demo database. If you place real files in the demo database, they will be deleted during future CODE-STAT software upgrade installations.

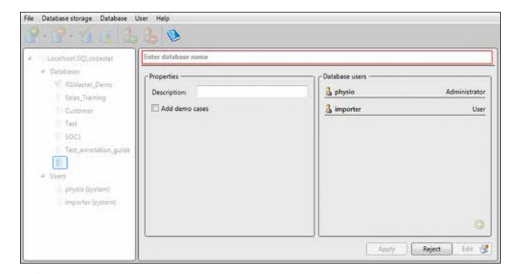
To create a new database, open CODE-STAT and log in. From the case list, go to Utilities → Database Utilities. Click on Add a new database or Edit an existing database configuration. Enter CODE-STAT administrator credentials (default name-physio, default password-control). Click **OK**.

From the CODE-STAT Database Utilities screen, click on the New Database icon 11.



Click on Create new database.

Enter database name in open box (don't use spaces between words; use underscore instead). Complete description, if desired and click apply.



Tip: If others need to be able to access the database for viewing cases or annotating from another desktop within your network, place the database on a network server where other computers have access.

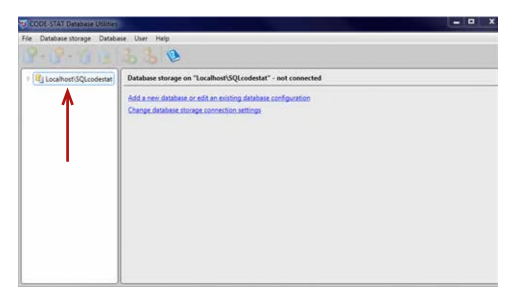
Once the new database is created, close the utility and open the new database by going to **File → Select Database** and select the new database from the dropdown menu. Continue to use the original demo username and password until you have set up new users.

NOTE: Files processed by CODE-STAT importer are imported to the database selected as default.

New Users

You can set up unique users and assign permission level. This is helpful if you have multiple users who will be accessing CODE-STAT software to view, edit, or annotate files.

To set up users, go to **Utilities** → **Database Utilities**. Double click **localhost** SQLcodestat.



Tip: Make sure users who will be responsible for editing and annotating cases have either a 'user' or 'administrator' access level.

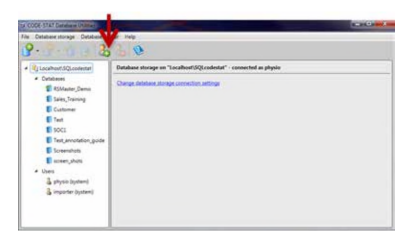
NOTE: You might want to create a common 'guest' account with an easy username and password. This allows providers to look at cases for educational purposes, but prevents them from changing or deleting the cases from the database.

Enter CODE-STAT administrator credentials.

Name: physio (default)

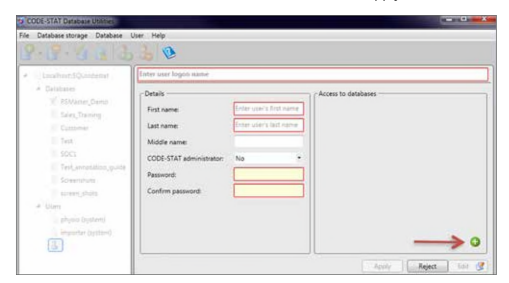
Password: control (default)





(Alternately, you can go to **User** → **Create user** on the toolbar.)

Fill in the red boxes, select Access to databases, then click Apply.



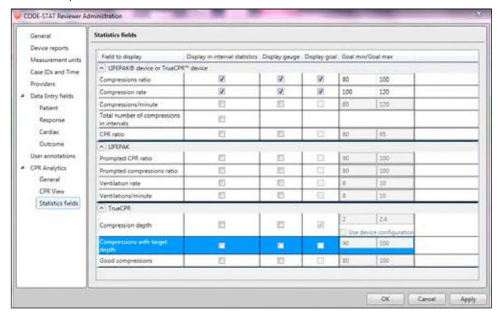
The new user's name will appear on the left hand side of the screen under Users.

CPR Performance Gauges

To set up CPR performance gauges (displayed on CPR report) for the first time, go to the Case List view → File → Administration → CPR Analytics → Statistics fields.

Set this up exactly as seen below:

- Compression ratio
- Compressions/minute



NOTE: Annotation of other dials will be covered in an advanced handbook.

NOTE: If you use continuous capnography and have determined your LIFEPAK 15 monitor/defibrillator has the capacity to record continuous waveform data in CODE-STAT (see page 2), you may also want to click ventilation rate and ventilations/minute since the program, in this case, uses an automatic ventilation detection algorithm.

| Case 1:22-cv-01378-MN-JLH | Document 175-2 1145 | Page 317 of 39 | 98 PageID #: |
|---------------------------|------------------------|----------------|--------------|
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |

Annotation Basics

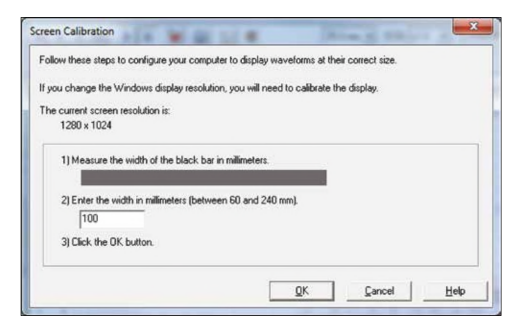
Setting Up Annotator Preferences

For training purposes, we suggest each annotator set up a copy of CODE-STAT data review software so they can follow along with this handbook.

Open any case in the database by double clicking on it.

Screen Calibration

If this is the first time opening a case in the database, a screen calibration box should appear. If not, go to View → Calibrate.

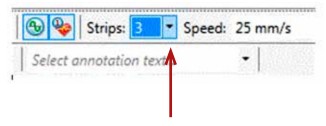


- 1. Measure the width of the black bar using a ruler.
- 2. Enter the measurement in mm.

NOTE: Calibrating the screen adjusts the display of waveforms so that the grid measurements are as close as possible to the actual measurements.

Settings

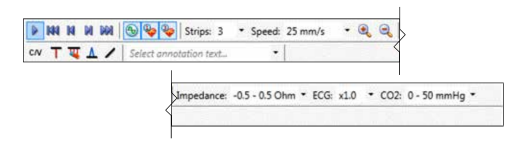
- Maximize the CODE-STAT program to cover the full screen.
- The case defaults to Landscape view. To change to Portrait view, go to View → Portrait on the toolbar.
- Select the number of strips visible on the screen by selecting 1-8 from the drop down menu.



 To customize settings for CPR Performance Gauges see Part 1: Introduction and setup.

NOTE: Choosing a larger number of strips will show more data on the screen but some of the channels (e.g., impedance, ECG) will overlap waveforms. Choose a lesser number of strips to keep the waveforms separated.

Annotation Tools



Playback Controls

| | Moves back one displayed page |
|----|---|
| | Moves forward one displayed page |
| N | Moves back one displayed strip (one row) |
| | Moves forward one displayed strip (one row) |
| | Plays the case in real time |
| 11 | Pauses a case playing in real time |

Channels and Waveform Displays

| (4) | Show or hide impedance signal waveform |
|-----------------------------|--|
| Q | Show or hide waveform from Channel 1 (e.g., ECG) |
| 2 | Show or hide waveform from Channel 2 (e.g., SpO ₂) |
| 3 | Show or hide waveform from Channel 3 (e.g., capnography) |
| Strips: 3 * | Select number of waveform strips displayed simultaneously |
| Speed: 25 mm/s * | Select sweep speed (horizontal time scale) of displayed waveform |
| વ્ વ | Increases or decreases sweepspeed |
| Impedance: -0.5 - 0.5 Ohm • | Increases or decreases size of impedance waveform |
| ECG: x1.0 | Select gain of ECG |
| CO2: 0 - 50 mmHg * | Select scale for viewing capnography waveform |

NOTE: When non-default values are selected, they are highlighted in yellow.

| Anr | ıotat | ions |
|-----|-------|------|
| | | |

C/V Shows or hides CPR events (compressions and ventilations)

on the waveform

Turns on/off compression editing mode

Turns on/off compression deleting mode

(multiple compressions simultaneously)

↑ Turns on/off ventilation editing mode

Turns on/off CPR break mode

Choose **annotations** from dropdown list then click on **waveform** to add desired annotation

Tip: To add custom events, go to Case list view. Go to File → Administration → User Annotations and enter your custom event in the bottom box and click ⊕.

Navigation Timeline Bars



The green navigation timeline represents the impedance signal over the length of the case. The highlighting frame indicates the currently displayed segment of the waveform.

Tip: To choose the ECG signal instead of the impedance signal, right-click the timeline and select **Channel 1 (ECG).** You can also view ETCO₂ or True CPR signals, if available.

Analysis
Shock
12-Lead

Device annotation

Compression period

ROSC period

Period excluded from the CPR report

U Current playback position

Tip: Click the markers to navigate to specific positions on the timeline.

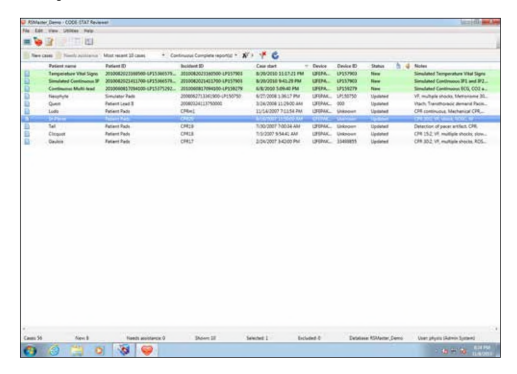
Suggested Basic Workflow

In order to implement a continuous quality improvement process based on data collected by LIFEPAK devices and reviewed through CODE-STAT software, we recommend the following basic workflow.



Opening A Case

To open a case, first open CODE-STAT data review software, select the database and log in. Double click on the case from the case list that needs annotation.



Annotating a Case

Here are the steps needed to annotate a case:

- 1. Open CODE-STAT data review software and log in to the database.
- 2. Open the case by double clicking on it.
- 3. Click on **Data Entry** tab **to enter patient's name and demographics** as needed.
- 4. Check that it is not a difficult case (more information below).
- 5. Mark "Start CPR Report" annotation.
- 6. Move through the case adding/deleting compressions and ventilations as needed.
- 7. Mark "ROSC" and "End ROSC" if found.
- 8. Verify "Stop CPR Report" annotation.
- 9. Close case and make "Closing Remarks," if desired.
- 10. Check your work.
- 11. Create and Print CPR Report.
- 12. Close case.

NOTE: Ventilations will be automatically annotated *only* when waveform capnography is visible.

NOTE: "Stop CPR Report" defaults to last compression. To change, delete annotation and re-annotate at desired location in case.

Steps 1-3 are self-explanatory.

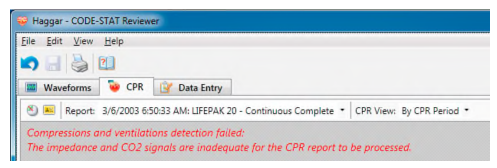
Step 4—Recognizing difficult cases.

Open the case. The impedance channel is visible over the ECG. Click [CN] to display automated chest compression (and ventilations if capnography waveform present).

Some cases are harder to annotate than others. It is recommended that you pass over the more difficult cases until you have had some practice. Look for the following to determine a difficult case:

For LIFEPAK 12 and certain 15 devices

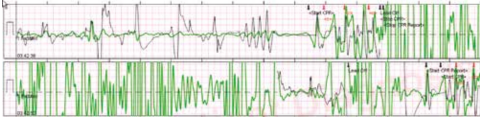
- No Impedance Signal Found. If there is no impedance signal, this means the defibrillator was not in PADDLES lead and therefore there is no impedance signal recorded. The "Show CPR Events" button or the Compression button
 - 👅 is inactive. A CPR Report 🕑 cannot be created without impedance data.



Example 4.1

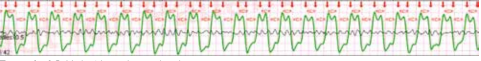
NOTE: The LIFEPAK 20/20e monitor/defibrillator and LIFEPAK 15 monitor/ defibrillator with (1) icon on the label located on the back of the device in the battery well, will record the impedance signal as long as defibrillation electrodes (pads) are applied to the patient. In these devices, you do not need to be in PAD-DLES LEAD in Channel 1 to record the impedance channel.

 Large amounts of artifact. Artifact of the impedance signal will make it difficult to accurately mark compressions.

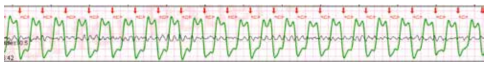


Example 4.2

• Variant impedance signal. If variant signals are present for only a brief period, this can easily be corrected by removing every other <c> marker (see example below). However, correcting long periods is a time-consuming effort.



Example 4.3: Variant impedance signal.



Example 4.4: Variant impedance signal corrected.

Tip: If the impedance signal height is too tall (out of range), simply decrease the size using the sizing tool Impedance: -0.5 - 0.5 Ohm + .

Step 5—Start CPR Report.

Turn on the "Show CPR Events button" C/V .

Turn on the "Compression editing button" T.

Determine where you would like to start measuring and add the first annotation, "Start CPR Report". The start of the report would typically coincide with the first sign of cardiac arrest. Sometimes the defibrillator has been powered on prior to the pads being connected and applied to the patient (as seen in example 5.1). When this is the case, place the annotation "Start CPR Report" at the first ECG waveform indicating cardiac arrest.

Note: "Start CPR Report" is different than "Start CPR."



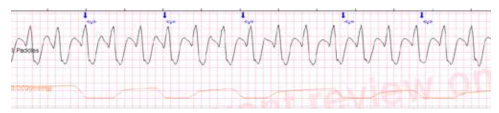
Step 6—Annotating compressions and ventilations.

A green impedance signal should be visible. Changes in the impedance signal due to compressions are recognized by the software, and are automatically annotated with a red arrow and <c> for compression. The software will place a blue arrow and <v> for ventilation if the capnography waveform is visible. The software will pick up 90-95% of the compressions and ventilations and annotate them automatically. Check the software's work, and add and delete compression and ventilation annotations as needed. Commonly, compressions may be missed at the beginning and end of cycles (note the first two compressions are not annotated correctly in example 6.1). Additionally, compressions may be missed when the signal pattern changes. Click To edit compressions. Hover over the compression with the mouse and click to add a compression annotation. Conversely, to remove an unwanted <c>, hover over it until the X appears and click to remove it. It is not imperative that the annotation is in an exact spot as long as there is a <c> for every compression.



Example 6.1: Missing compressions

To edit ventilations click on \triangle and proceed as with compressions.

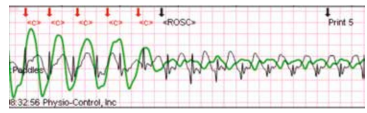


Example 6.2: Ventilation annotation

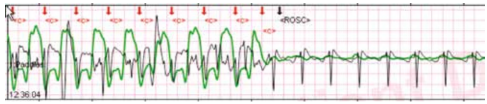
Tip: Many variables go into the makeup of the impedance signal. It is not reasonable to interpret the height of the signal to be an indicator of depth or quality of the provider's compression. The impedance signal can only give you rate and number of compressions.

Step 7—Marking ROSC (Return of Spontaneous Circulation) and End ROSC.

When return of spontaneous circulation is obtained, use the drop down menu to select **ROSC** and mark the annotation.



Example 7.1: ROSC marked



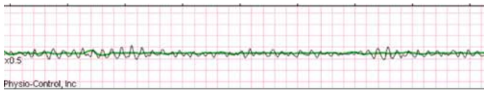
Example 7.2: ROSC marked

Marking ROSC will tell the software that the following time frame should not have compressions. This makes the CPR Report accurate.

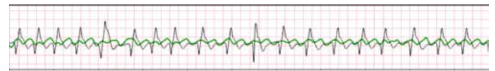
Looking for the following information provided in the case file or other patient documentation records can assist in determining ROSC. The more that are present, the higher the probability of ROSC.

- Providers stopped CPR
- A narrow QRS complex
- CPR is not started again in a reasonable amount of time
- ETCO2 rises
- A blood pressure is documented
- Verbalization of ROSC (if voice recording is available)
- A provider event marks ROSC or charts ROSC on the medical report

Because blood is a great electrical conductor, a rhythmic low amplitude pattern to the impedance signal may be present, usually around the T wave on the ECG (see examples 7.3 and 7.4 below).

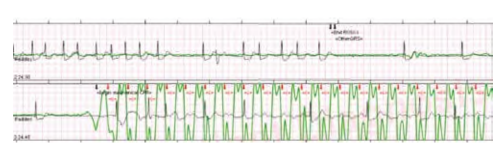


Example 7.3: Notice the impedance line is very flat. No blood flow.



Example 7.4: Notice the impedance signal rhythmic pattern near the T wave, suggestive of blood flow associated with ROSC.

If ROSC is lost, generally compressions will begin again. Attempt to determine where ROSC was lost and use the drop down menu to select "End ROSC" and mark the annotation.



Example 7.5: Mark where "End ROSC" is most likely. Sometimes it will be right at the start of CPR or, like in this case, where the rhythm deteriorates before the provider notices and takes action.

Step 8—Verify "Stop CPR Report" annotation.

This annotation tells the software the point at which performance measurements should stop (if ROSC is not annotated) and is automatically placed at the last compression. This avoids inaccurate statistics if the device is left on after resuscitation measures have ceased. (see example 8.1).

If you want to edit this annotation you must first delete the original annotation, determine a different end point and select STOP CPR report from the dropdown box.

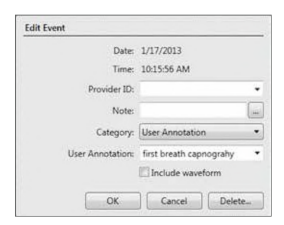
NOTE: The default setting can be changed in File → Administration → General (under CPR Analytics).



Example 8.1: CPR stopped. Resuscitation ceased and a "Stop CPR Report" is annotated at the last compression.

Step 9—Closing remarks and Special Annotations.

To add comments or special annotations, left click on the waveform to bring up the **Edit Event** box shown below. Select a category from the pull down menu (User annotation, Medication, Procedure CPR annotation), then select from the various choices in the next drop down window.



Tip: If it is difficult to annotate ROSC, a note may be added for future reference as to the reason ROSC was annotated where it was.

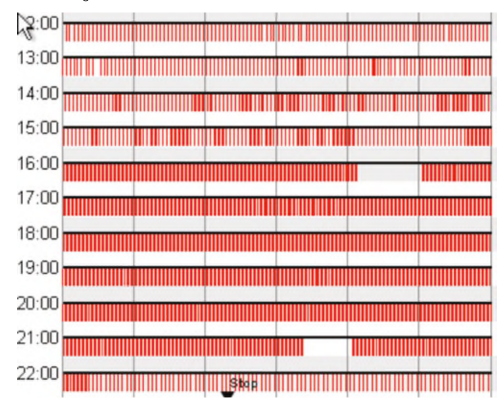
Step 10—Checking Over the Work.

Checking over the work is important to assure accuracy. Review for the following:

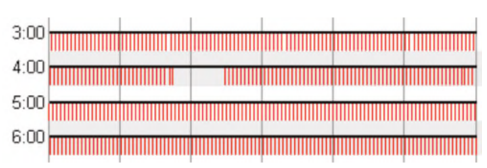
- Is there a Start CPR Report annotation?
- If there is ROSC, is it marked?
- How about loss of ROSC?
- Was ROSC regained?
- If the case was terminated, is there a Stop CPR Report annotation?

In addition, review the case for correct annotation of compressions by creating a CPR Report (Example 10.1).

- Look for large gaps, as well as areas where compressions appear denser (darker). Click anywhere on the CPR Report to be transported directly to that part of the case. Delete incorrect <c> annotations. Add missing <c> annotations.
- If capnography waveform is visible, delete incorrect <v> annotations. Add missing <v> annotations.

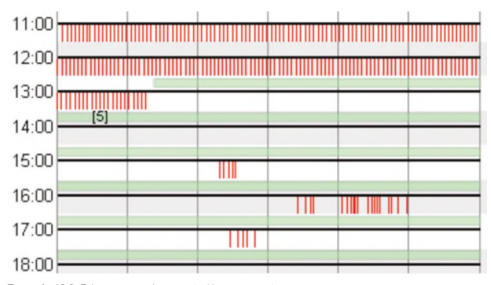


Example 10.1: Notice the denser (darker) areas in minute 13 forward. This is caused by twice the number of compression annotations. Click on this area to go directly to that area and clean it up.



Example 10.2: Notice how the compression annotations look clean and rhythmic. There are no areas that are denser than others. This is a sign of good CPR performance and an accurate record.

 Look for and remove false compressions (generally due to movement of the patient) in the ROSC area if applicable (Example 10.3). Although compressions located in the ROSC area do not alter performance measures, they can look confusing on the CPR Report.

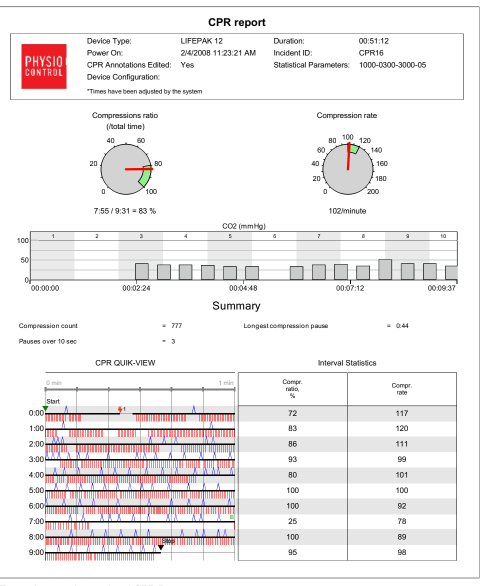


Example 10.3: False compressions created by movement.

Step 11—Creating and Printing the CPR Report.

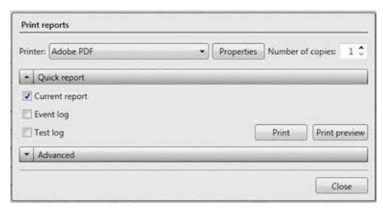
Once the work is complete, re-create a CPR Report by clicking the PR tab.

When the CPR Report appears, be sure to click to display the performance dials.



Example 11.1: A completed CPR Report.

The report can be printed to a physical printer or a PDF file. There are two ways. From the report view, click on the printer icon → This brings up another screen. Select the printer and click → or go to File → Print → Case Reports. Current report is the default. Click Print.



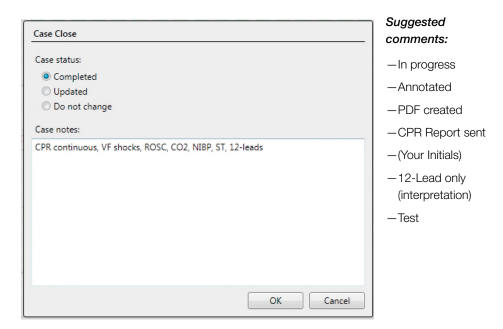
Example 11.2: Printing the CPR Report.

Tip: If you want an electronic version of the report, change the Printer selection to Adobe PDF and save to your files.

Step 12—Closing a Case—Closing Remarks.

To close a case and return to the case list click on the toolbar.

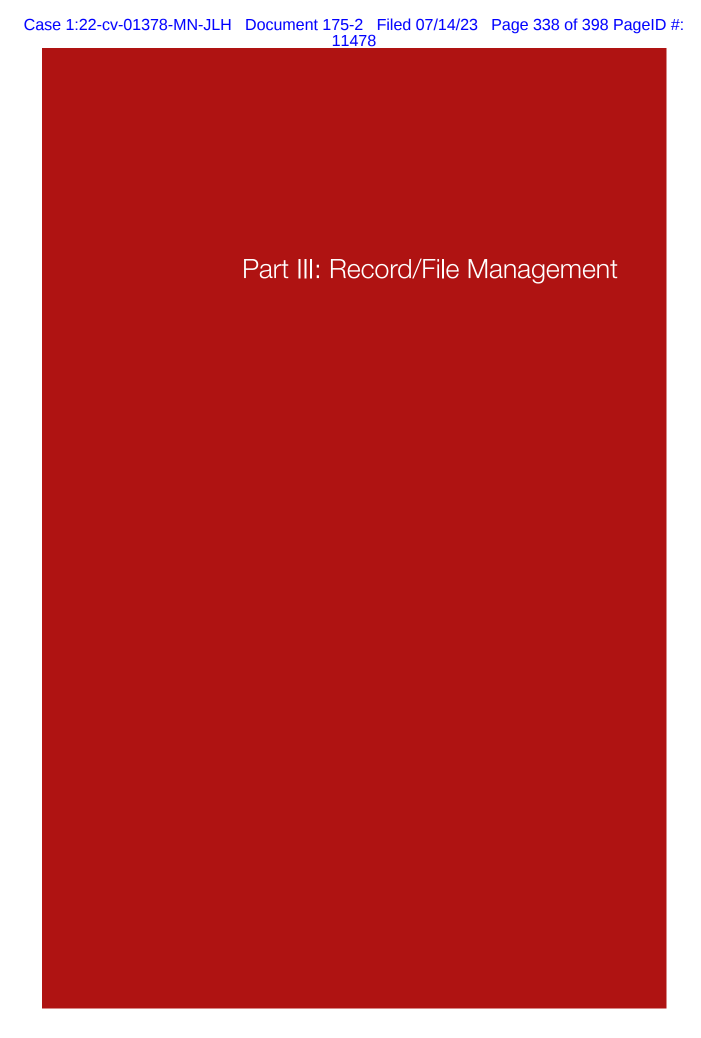
When closing out of a case, take the opportunity to document what has been completed. This is helpful because any remarks will show up on the case list, making it easier to locate the case later.



The Case Close dialog box appears every time you close a case. Before you can close the case, you must indicate its current status:

• Completed—If you finished all activities related to the report.

NOTE: If you don't update or complete the status for the case, it will remain as NEW CASE in the case list.



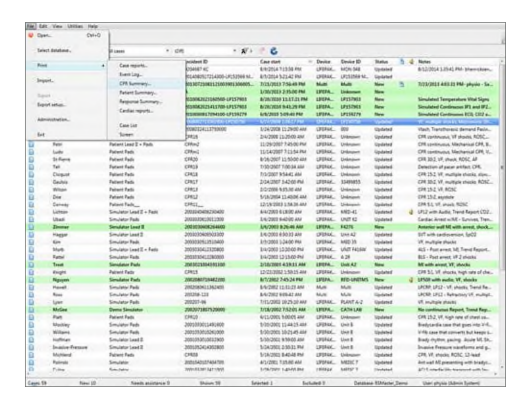
Creating Summary CPR Reports

Creating Multi-Case Reports

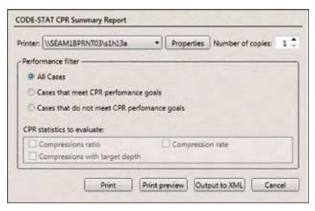
Multi-case reports are helpful for trending, monthly meeting updates and yearly reports.

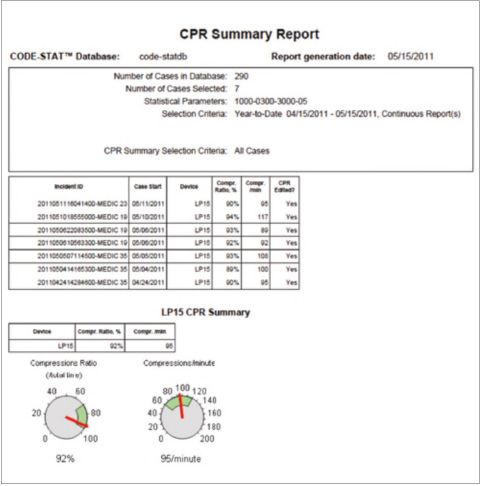
To create monthly or custom reports that look at periods of time and provide information for all the cardiac arrests during that period:

- 1. Open CODE-STAT and login to the appropriate database.
- 2. Filter the case list to show the reports you would like to include. The report will run on all of the cases displayed in the report list.
- 3. Click File \rightarrow Print \rightarrow CPR Summary.



4. Change the performance filter, if desired (default is set to all cases) and click **Print**.





Example: Monthly Report.

Importing and Exporting Cases

Importing Cases into CODE-STAT Data Review Software

You can automatically import cases into CODE-STAT via:

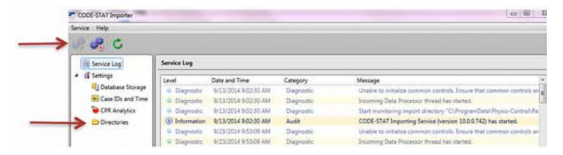
- LIFENET® Connect
- DT EXPRESS™ 6.0 Data Transfer Software

LIFENET Connect is used to send files to CODE-STAT data review software directly from LIFEPAK devices in the field. This is typically part of a larger implementation. Your LIFENET implementation specialist will assist with this setup.

DT EXPRESS is a stand-alone software program used to download cases from the LIFEPAK devices. It is also used to send data to CODE-STAT data review software for storage/review/annotation. CODE-STAT 10 no longer has local download capabilities included in the program (like CODE-STAT 9 did). Instead CODE-STAT 10 can link to DT EXPRESS 6.0 if they are installed on the same computer. To do this, DT EXPRESS 6.0 needs to be configured in order to move files into CODE-STAT 10 automatically. One way to do this is to setup an automatic exporter in to DT EXPRESS 6.0 that moves the file into the importer folder automatically.

Automatic Import of Files into CODE-STAT Database

To import files automatically via LIFENET Connect or DT EXPRESS, make sure CODE-STAT Import Service is running. To check, go to the Start menu on your computer and click all programs → Physio-Control → Post Event Review → CODE-STAT 10 → CODE-STAT importer. If sis displayed, the Import Service is already running. If 🥒 is displayed, click on this icon to start the Import Service.



Now CODE-STAT Import Service is running and files will automatically be imported.

To configure DT EXPRESS 6.0 data transfer software to automatically move files to CODE-STAT:

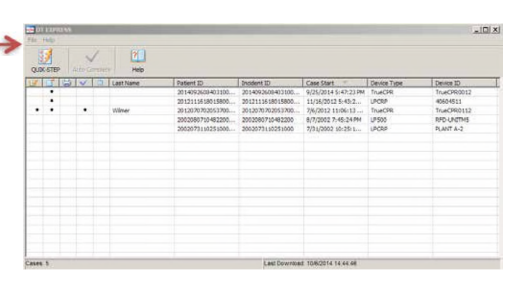
- 1. Click on Directories.
- 2. Copy "Import files from" path this will be used to configure an exporter in DT EXPRESS.



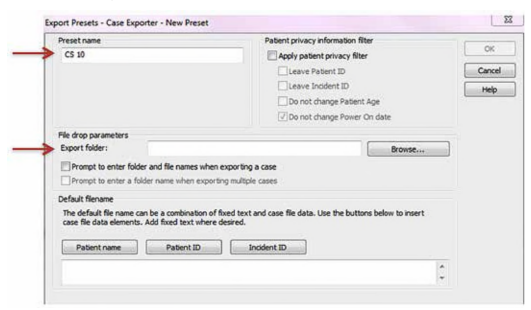
3. Open the DT EXPRESS software by clicking on the toolbar in CODE-STAT data review software.



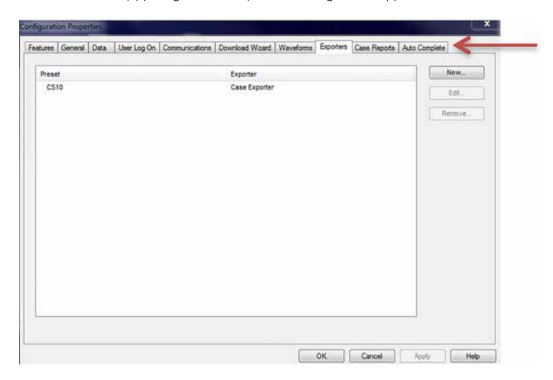
NOTE: If is disabled, you will need to install DT EXPRESS 6.0 data transfer software on the same computer as your CODE-STAT 10 data review software.



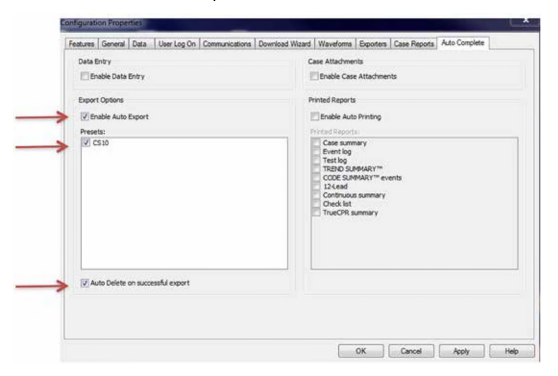
- Select File → Configuration → Exporters tab → New → Case Exporter → OK.
- 5. Fill in preset name (e.g., CS 10).
- 6. Paste "Importer files from" location (copied in step 2) into Export folder field.



7. Click **OK** (upper right of screen). The following screen appears.



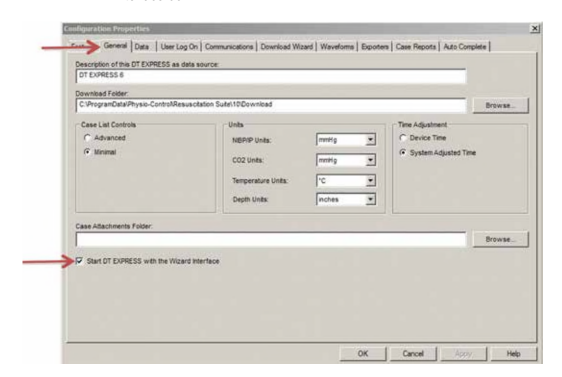
8. Select Auto Complete tab.



- 9. Check Enable Auto Export under Export Options.
- 10. Select Presets (e.g. CS10).
- 11. If you want to delete local copy, select Auto Delete on successful export (recommended).

Once configured, files will automatically move into your default CODE-STAT database.

12. To have the Quick Step screen come up automatically in the future, go to the General tab and check Start DT EXPRESS with the Wizard interface box.



13. Click OK

Now the screen below will appear when you open DT EXPRESS. Click on NEXT and follow directions.



Manual Import of Files into CODE-STAT Database

If you already have a Physio Case Object (PCO) file (i.e., delivered via email), you can manually import that file into CODE-STAT software. To do so, go to the case list and then click File -> Import and then navigate to where you saved the PCO file on your computer. Refresh the case list and the imported case will appear in the list.

You can also manually download a case into CODE-STAT from a LIFEPAK device using the above DT EXPRESS 6.0 configuration.

NOTE: You cannot view ECG waveforms on the screen or playback audio reports in DT EXPRESS software. To do this, you need to export a case to CODE-STAT software.

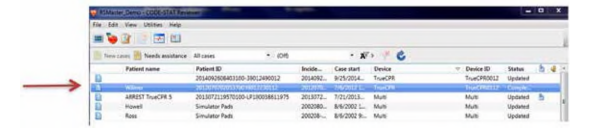
NOTE: For detailed instructions regarding the use of DT EXPRESS data transfer software, please refer to $Help \rightarrow DT$ EXPRESS Help.

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TrueCPR™ Coaching Device

Viewing TrueCPR Device Data

Files that contain data from the TrueCPR coaching device and have been imported into CODE-STAT 10 data review software can be viewed by double clicking on the file in the case list/reviewer view.



The following data can be displayed:

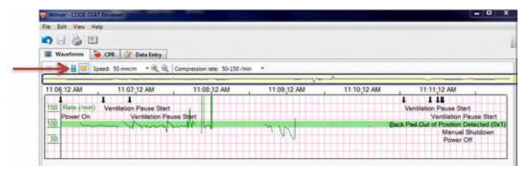
- Compression depth diagram
- Compression rate diagram

Target zones for depth and rate are indicated by thick solid lines.

NOTE: These colors may be changed in Graphic Schemes.



Click **!** on the tool bar to show/hide depth peaks diagram.



Example of depth waveform hidden from view. Only rate information is visible.

Click **!** again to make depth information visible again.

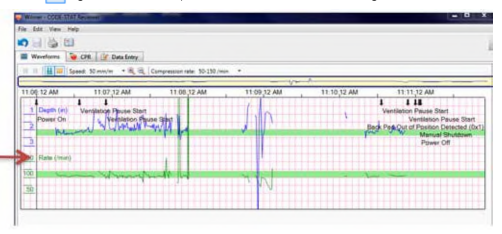


Click on the tool bar to show/hide compression rate diagram.

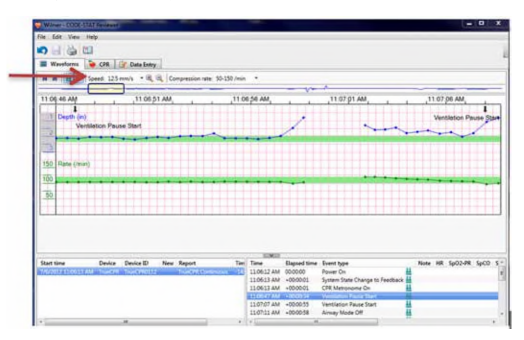


Example of compression rate hidden from view. Only compression depth information is visible.

Click again to make compression rate information visible again.



Increasing the speed will spread the data out so it is easier to view compressions individually.



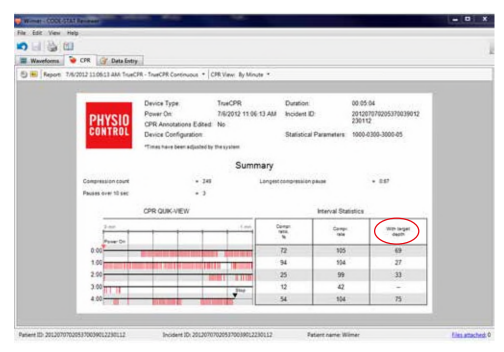
NOTE: Ventilation Pause Start is device annotation in the TrueCPR device when it is set to the No Airway mode (30:2).

CPR QUIK-VIEW™ Data Review Program Summary with TrueCPR Data

To view a CPR QUIK-VIEW summary with TrueCPR data:

Click properties on the toolbar.



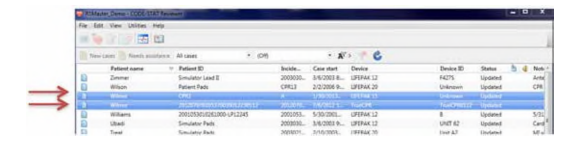


Example: CPR QUIK summary with TrueCPR data.

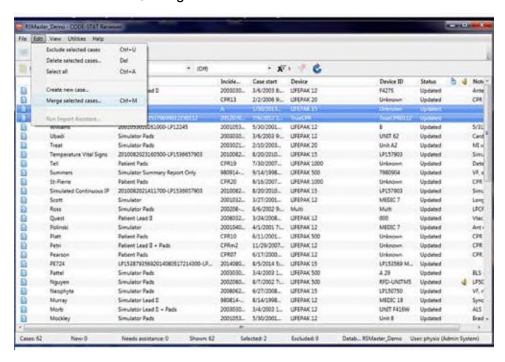
Combining Patient Data from TrueCPR and LIFEPAK Devices

To combine patient data from the TrueCPR device with the patient data from the LIFEPAK monitor/defibrillator by merging the two cases:

1. Highlight the two cases you want to merge.

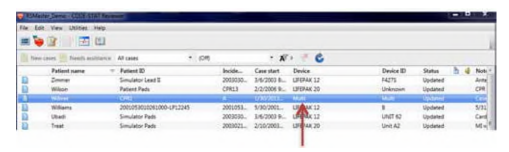


Go to Edit → Merge select cases.



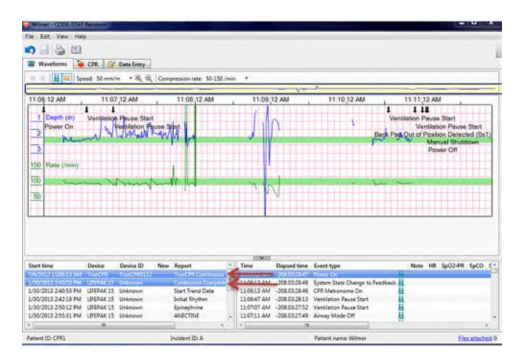
3. Follow the Wizard that will guide you through the steps of merging the two selected cases into one case.

Once the cases are merged, you will see one case with Multi under Device column.

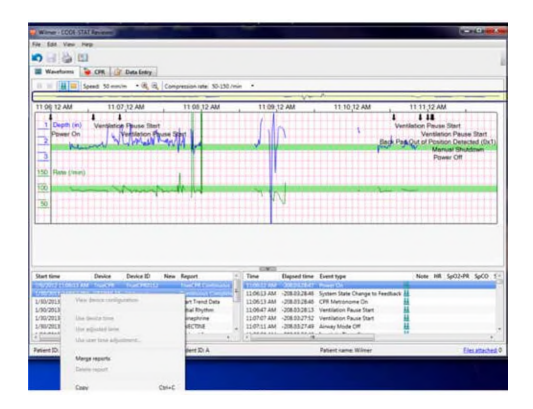


Viewing Combined LIFEPAK monitor/defibrillator and TrueCPR Device Data

To view the LIFEPAK monitor/defibrillator and TrueCPR device data together in the continuous viewer and the CPR QUIK-VIEW summary report, you will need to merge the continuous reports located in the report list from the two devices. Double click on the merged case to open the viewer and go to the report list.



- 1. Highlight the continuous report and the TrueCPR continuous report.
- 2. Right click on highlighted cases and select Merge reports.



Now the TrueCPR data is combined with the continuous ECG data in a new report displayed in the Report list as Multi under Device column.



NOTE: If the compression waveform and the TrueCPR compressions (blue dots) don't line up, use Shift TrueCPR arrows to align.

NOTE: CODE-STAT 10 software cannot combine patient data between LIFEPAK devices (e.g., LIFEPAK 15 monitor/defibrillator to LIFEPAK 1000 AED) into one view.

NOTE: More data can be included by choosing options under the CPR Analytics tab of the Administration window.

For further information please contact your local Physio-Control representative or visit our website at www.physio-control.com.



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Protective Order

IN THE UNITED STATES DISTRICT COURT FOR THE DISTRICT OF DELAWARE

| APPLE INC., |) |
|---|-------------------------------------|
| Plaintiff, |))) C.A. No. 22-1377-MN-JLH |
| v. |)) JURY TRIAL DEMANDED |
| MASIMO CORPORATION and SOUND UNITED, LLC, |)) |
| Defendants. |) |
| MASIMO CORPORATION, | <u>)</u> |
| Counter-Claimant, |) |
| v. |) |
| APPLE INC., |) |
| Counter-Defendant. |) |
| APPLE INC., | |
| Plaintiff, |) |
| v. |) C.A. No. 22-1378-MN-JLH |
| MASIMO CORPORATION and SOUND UNITED, LLC, |) JURY TRIAL DEMANDED |
| Defendants. |) |
| MASIMO CORPORATION and CERCACOR LABORATORIES, INC., |) |
| Counter-Claimants, |) |
| v. | ,)) |
| APPLE INC., | ,)) |
| Counter-Defendant. | ,) |

AGREED PROTECTIVE ORDER REGARDING THE DISCLOSURE AND USE OF DISCOVERY MATERIAL

Plaintiff and Counter-Defendant Apple Inc. ("Plaintiff"), Defendants and Counter-Claimants Masimo Corporation and Sound United, LLC and Counter-Claimant Cercacor Laboratories, Inc. (together, "Masimo") anticipate that documents, testimony, or information containing or reflecting confidential, proprietary, trade secret, and/or commercially sensitive information are likely to be disclosed or produced during the course of discovery, initial disclosures, and supplemental disclosures in these cases and request that the Court enter this Order setting forth the conditions for treating, obtaining, and using such information.

Pursuant to Rule 26(c) of the Federal Rules of Civil Procedure, the Court finds good cause for the following Agreed Protective Order Regarding the Disclosure and Use of Discovery Material ("Order" or "Protective Order").

1. PURPOSES AND LIMITATIONS

- (a) Protected Material designated under the terms of this Protective Order shall be used by a Receiving Party solely for these cases, and shall not be used directly or indirectly for any other purpose whatsoever.
- (b) The Parties acknowledge that this Order does not confer blanket protections on all disclosures during discovery, or in the course of making initial or supplemental disclosures under Rule 26(a). Designations under this Order shall be made with care and shall not be made absent a good faith belief that the designated material satisfies the criteria set forth below. If it comes to a Producing Party's attention that designated material does not qualify for protection at all, or does not qualify for the level of protection initially asserted, the Producing Party must promptly notify all other Parties that it is withdrawing or changing the designation.

(c) Other Proceedings. By entering this order and limiting the disclosure of information in these cases, the Court does not intend to preclude another court from finding that information may be relevant and subject to disclosure in another case. Any person or party subject to this order who becomes subject to a request or motion that would require disclosure of another party's information designated "CONFIDENTIAL," "CONFIDENTIAL - ATTORNEYS' EYES ONLY," or "CONFIDENTIAL - OUTSIDE ATTORNEYS' EYES ONLY - SOURCE CODE," pursuant to this Order shall promptly notify that party of the request or motion so that the party may have an opportunity to appear and be heard on whether that information should be disclosed.

2. **DEFINITIONS**

- (a) "Affiliate" means any corporation, company, or other business entity over which a Party has the power to direct or cause the direction of the management, policies, or legal actions through: (1) at least 50% ownership of voting securities; or (2) contract; or (3) other means.
- (b) "Discovery Material" means all items or information, including from any non-party, regardless of the medium or manner generated, stored, or maintained (including, among other things, testimony, transcripts, or tangible things) that are produced, disclosed, or generated in connection with discovery or Rule 26(a) disclosures in these cases.
- (c) "Outside Counsel" means (i) outside counsel who appear on the pleadings as counsel for a Party and (ii) partners, associates, and staff of such counsel to whom it is reasonably necessary to disclose the information for this litigation.
- (d) "Patents-in-suit" means U.S. Patent Nos. D735,131, D883,279, D947,842, D962,936, 10,076,257, 10,627,783, 10,942,491, 10,987,054, 11,106,352, 11,474,483, 10,912,501, 10,912,502, 10,945,648, 10,687,743, 10,687,745, 10,722,159, 7,761,127, 8,190,223, 10,736,507,

and 10,984,911 and any other patent asserted in these cases, as well as any related patents, patent applications, provisional patent applications, continuations, and/or divisionals.

- (e) "Party" means any party to these cases, including all of its officers, directors, employees, consultants, vendors, retained experts, and outside counsel and their support staffs.
- (f) "Producing Party" means any Party or non-party that discloses or produces any Discovery Material in these cases.
- (g) "Protected Material" means any Discovery Material that is designated as "CONFIDENTIAL," "CONFIDENTIAL ATTORNEYS' EYES ONLY," or "CONFIDENTIAL OUTSIDE ATTORNEYS' EYES ONLY SOURCE CODE," as provided for in this Order. Protected Material shall not include: (i) advertising materials that have been actually published or publicly disseminated; and (ii) materials that show on their face they have been disseminated to the public.
- (h) "Receiving Party" means any Party who receives Discovery Material from a Producing Party.
- (i) "Source Code" means computer code, scripts, assembly, binaries, object code, source code listings (e.g., file names and path structure), descriptions of source code (e.g., descriptions of declarations, functions, and parameters), object code listings and descriptions of object code, Hardware Description Language (HDL) or Register Transfer Level (RTL) files that describe the hardware design of any ASIC or other chip, and native Computer Aided Design (CAD) files that describe the hardware design of any component, the disclosure of which to another Party or non-party is likely to cause harm or competitive disadvantage to the Producing Party. To avoid any doubt, still images of CAD files are not Source Code and will not be subject to the

disclosure and review restrictions in Section 11. Still images of CAD files may be designated as "CONFIDENTIAL" or "CONFIDENTIAL - ATTORNEYS' EYES ONLY," as provided for in this Order.

3. <u>COMPUTATION OF TIME</u>

The computation of any period of time prescribed or allowed by this Order shall be governed by the provisions for computing time set forth in Federal Rules of Civil Procedure 6.

4. **SCOPE**

- (a) The protections conferred by this Order cover not only Discovery Material governed by this Order as addressed herein, but also any information copied or extracted therefrom, as well as all copies, excerpts, summaries, or compilations thereof, plus testimony, conversations, or presentations by Parties or their counsel in court or in other settings that might reveal Protected Material.
- (b) Nothing in this Protective Order shall prevent or restrict a Producing Party's own disclosure or use of its own Protected Material for any purpose, and nothing in this Order shall preclude any Producing Party from showing its Protected Material to an individual who prepared the Protected Material.
- (c) Nothing in this Order shall be construed to prejudice any Party's right to use any Protected Material with the consent of the Producing Party or by order of the Court.
- (d) This Order is without prejudice to the right of any Party to seek further or additional protection of any Discovery Material or to modify this Order in any way, including, without limitation, an order that certain matter not be produced at all.

(e) Any use of Protected Material at trial shall be governed by the orders of the trial judge and other applicable authorities. This Order does not govern the use of Protected Material at trial.

5. **DURATION**

Even after the termination of these cases, the confidentiality obligations imposed by this Order shall remain in effect until a Producing Party agrees otherwise in writing or a court order otherwise directs.

6. <u>ACCESS TO AND USE OF PROTECTED MATERIAL</u>

- (a) <u>Basic Principles</u>. All Protected Material shall be used solely for these cases or any related appellate proceedings, and not for any other purpose whatsoever, including without limitation, any other litigation, patent prosecution or acquisition, patent reexamination or reissue proceedings, or any business or competitive purpose or function. Protected Material shall not be distributed, disclosed, or made available to anyone except as expressly provided in this Order.
- Outside Counsel and any person associated with a Party who receives a Producing Party's material designated "CONFIDENTIAL ATTORNEYS' EYES ONLY" or "CONFIDENTIAL ATTORNEYS' EYES ONLY SOURCE CODE" under this Protective Order or who has access to, accesses, or otherwise learns of, in whole or in part, said material designated "CONFIDENTIAL ATTORNEYS' EYES ONLY" or "CONFIDENTIAL ATTORNEYS' EYES ONLY" or "CONFIDENTIAL ATTORNEYS' EYES ONLY SOURCE CODE" under this Protective Order shall not prepare, prosecute, supervise, advise, counsel, or assist in the preparation or prosecution of any patent application seeking a patent on behalf of the Receiving Party or its acquirer, successor, predecessor, or Affiliate in the field of non-invasive monitoring and/or consumer wearables (generally or as

described in any patent in suit) during the pendency of this Action and for two years after final termination of this action, including all appeals. To avoid any doubt, "prosecution" as used in this section does not include representing or advising a Party before a domestic or foreign agency in connection with a reissue, ex parte reexamination, covered business method review, inter partes review, opposition, cancelation, or similar proceeding; though in connection with any such foreign or domestic agency proceeding involving the patents-in-suit, any attorney who has access to, accesses, obtains, receives, or otherwise learns, in whole or in part, any other Party's "CONFIDENTIAL – ATTORNEYS' EYES ONLY" or "CONFIDENTIAL – ATTORNEYS' EYES ONLY – SOURCE CODE" shall not: (i) participate in the preparation, prosecution, supervision, advice, counsel, or assistance of any amended claims; (ii) reveal a Producing Party's Protected Material to any prosecuting reexamination counsel or agent; or (iii) use a Producing Party's Protected Material for any purpose not permitted by Section 1.

(c) <u>Secure Storage</u>, <u>No Export</u>. Protected Material must be stored and maintained by a Receiving Party at a location in the United States and in a secure manner that ensures that access is limited to the persons authorized under this Order. To ensure compliance with applicable United States Export Administration Regulations, Protected Material may not be exported outside the United States or released to any foreign national, even if within the United States. This applies to such information regardless of whether it is in the form of a stand-alone document or as an exhibit, attachment, or appendix to anything, including but not limited to briefs, reports, letters to counsel, discovery responses, or court filings—whether drafts or final versions. Foreign nationals shall not include the Parties' Outside Counsel who reside in the United States, agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A, and who are identified in writing to the Producing Party. However, the Parties' Outside Counsel

may access briefs, reports, letters to counsel, discovery responses, and court filings (including drafts) that contain Protected Material for purposes of working on these cases while traveling temporarily outside the United States, exclusive of any exhibits or appendices that attach or substantially reproduce or summarize documents, data, or testimony that have been designated by any other party as Protected Material. The Parties will use their best efforts to minimize the amount of Protected Materials in those documents (including without limitation by redacting references to Protected Materials that are not necessary for the work performed outside of the United States) to help ensure the security of the Parties' Protected Materials. Also, if this case eventually requires depositions or experts located outside the United States, the parties will revisit this issue and attempt to agree about exporting specific materials to the extent necessary. The Parties agree that neither Party waives the right to seek amendment of this Protective Order by the Court, following a meet and confer, if other circumstances concerning exportation arise in this case.

- (d) <u>Legal Advice Based on Protected Material</u>. Nothing in this Protective Order shall be construed to prevent counsel from advising their clients with respect to these cases based in whole or in part upon Protected Materials, provided counsel does not disclose the Protected Material itself except as provided in this Order.
- (e) <u>Limitations</u>. Nothing in this Order shall restrict in any way a Producing Party's use or disclosure of its own Protected Material. Nothing in this Order shall restrict in any way the use or disclosure of Discovery Material by a Receiving Party: (i) that is or has become publicly known through no fault of the Receiving Party; (ii) that is lawfully acquired by or known to the Receiving Party independent of the Producing Party; (iii) previously produced, disclosed and/or provided by the Producing Party to the Receiving Party or a non-party without an

obligation of confidentiality and not by inadvertence or mistake; (iv) with the consent of the Producing Party; or (v) pursuant to order of the Court.

7. <u>DESIGNATING PROTECTED MATERIAL</u>

- (a) <u>Available Designations</u>. Any Producing Party may designate Discovery Material with any of the following designations, provided that it meets the requirements for such designations as provided for herein: "CONFIDENTIAL," "CONFIDENTIAL ATTORNEYS' EYES ONLY," or "CONFIDENTIAL OUTSIDE ATTORNEYS' EYES ONLY SOURCE CODE."
- (b) Written Discovery and Documents and Tangible Things. Written discovery, documents (which include "electronically stored information," as that phrase is used in Federal Rule of Procedure 34), and tangible things that meet the requirements for the confidentiality designations listed in Section 7(a) may be so designated by placing the appropriate designation on every page of the written material prior to production. For digital files being produced, the Producing Party may mark each viewable page or image with the appropriate designation, and mark the medium, container, and/or communication in which the digital files were contained. In the event that original documents are produced for inspection, the original documents shall be presumed "CONFIDENTIAL ATTORNEYS' EYES ONLY" during the inspection and re-designated, as appropriate during the copying process.
- (c) Native Files. Where electronic files and documents are produced in native electronic format, such electronic files and documents shall be designated for protection under this Order by appending to the file names or designators information indicating whether the file contains "CONFIDENTIAL," "CONFIDENTIAL ATTORNEYS' EYES ONLY," or "CONFIDENTIAL OUTSIDE ATTORNEYS' EYES ONLY SOURCE CODE," material, or

shall use any other reasonable method for so designating Protected Materials produced in electronic format. When electronic files or documents are printed for use at deposition, in a court proceeding, or for provision in printed form to an expert or consultant pre-approved pursuant to Section 12, the party printing the electronic files or documents shall affix a legend to the printed document corresponding to the designation of the Producing Party and including the production number and designation associated with the native file. The parties reserve the right to object to the use of any image format version of a document produced in native format to the extent any information has been altered.

Depositions and Testimony. Parties or testifying persons or entities may (d) designate depositions and other testimony with the appropriate designation by indicating on the record at the time the testimony is given or by sending written notice of how portions of the transcript of the testimony are designated within fifteen (15) days of receipt of the transcript of the testimony. If no indication on the record is made, all information disclosed during a deposition shall be deemed "CONFIDENTIAL - ATTORNEYS' EYES ONLY" until the time within which it may be appropriately designated as provided for herein has passed. Any Protected Material that is used in the taking of a deposition shall remain subject to the provisions of this Protective Order, along with the transcript pages of the deposition testimony dealing with such Protected Material. In such cases the court reporter shall be informed of this Protective Order and shall be required to operate in a manner consistent with this Protective Order. In the event the deposition is videotaped, the original and all copies of the videotape shall be marked by the video technician to indicate that the contents of the videotape are subject to this Protective Order, substantially along the lines of "This videotape contains confidential testimony used in this case and is not to be viewed or the contents thereof to be displayed or revealed except pursuant to the terms of the operative

Protective Order in this matter or pursuant to written stipulation of the parties." Counsel for any Producing Party shall have the right to exclude from oral depositions, other than the deponent, deponent's counsel, the reporter and videographer (if any), any person who is not authorized by this Protective Order to receive or access Protected Material based on the designation of such Protected Material. Such right of exclusion shall be applicable only during periods of examination or testimony regarding such Protected Material.

8. <u>DISCOVERY MATERIAL DESIGNATED AS "CONFIDENTIAL"</u>

- (a) A Producing Party may designate Discovery Material as "CONFIDENTIAL" if it contains or reflects confidential, proprietary, and/or commercially sensitive information.
- (b) Unless otherwise ordered by the Court, Discovery Material designated as "CONFIDENTIAL" may be disclosed only to the following:
- (i) The Receiving Party's Outside Counsel, such counsel's immediate paralegals and staff, and any copying or clerical litigation support services working at the direction of such counsel, paralegals, and staff;
- (ii) Officers or employees of the Receiving Party, who may be, but need not be, in-house counsel for the Receiving Party, as well as their immediate paralegals and staff, to whom disclosure is reasonably necessary for this case, provided that each such person has agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A;
- (iii) Any outside expert or consultant retained by the Receiving Party to assist in these cases, provided that disclosure is only to the extent necessary to perform such work; and provided that: (a) such expert or consultant has agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A; (b) such expert or consultant is not a current

officer, director, or employee of a Party or of a competitor of a Party, nor anticipated at the time of retention to become an officer, director or employee of a Party or of a competitor of a Party; (c) such expert or consultant accesses the materials in the United States only, and does not transport them to or access them from any foreign jurisdiction (however, to avoid doubt, such expert or consultant may access reports (including drafts) that contain the materials for purposes of working on these cases while traveling temporarily outside the United States); and (d) no unresolved objections to such disclosure exist after proper notice has been given to all Parties as set forth in Section 12 below;

- (iv) Witnesses at depositions or hearings in these cases and the witnesses' counsel, provided however that the disclosure shall only be made to: (1) a witness who is an employee of the Producing Party, or identified on the document as an author, addressee, or recipient of the material in question, or if there are other indicia (such as from metadata, cover emails, or other records of distribution) that the witness has seen or had access to the document previously; or (2) a witness who has been designated to testify on behalf of the Producing Party on the subject matter of the material in question, provided however that the Protected Material shown to such a witness shall be limited to Protected Material of the Producing Party;
- (v) Court reporters, stenographers and videographers retained to record testimony taken in these cases, and their staff;
 - (vi) The Court, jury, and court personnel;
- (vii) Graphics, translation, design, trial consulting personnel, and/or other professional vendors, having first agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A;

- (viii) Mock jurors having first agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A.
- (ix) Any mediator who is assigned to hear these matters, and his or her staff, subject to their agreement to maintain confidentiality to the same degree as required by this Protective Order; and
- (x) Any other person with the prior written consent of the Producing Party.

9. <u>DISCOVERY MATERIAL DESIGNATED AS "CONFIDENTIAL – ATTORNEYS" EYES ONLY"</u>

- (a) A Producing **Party** designate Discovery Material may as "CONFIDENTIAL – ATTORNEYS' EYES ONLY" if it contains or reflects information that is extremely confidential and/or sensitive in nature and the Producing Party reasonably believes that the disclosure of such Discovery Material is likely to cause harm or significant competitive disadvantage to the Producing Party. The Parties agree that the following information, if nonpublic, shall be presumed to merit the "CONFIDENTIAL - ATTORNEYS' EYES ONLY" trade secrets, pricing information, financial data, sales information, sales or marketing forecasts or plans, business plans, sales or marketing strategy, product development information, engineering documents, testing documents, employee information, and other nonpublic information of similar competitive and business sensitivity.
- (b) Unless otherwise ordered by the Court, Discovery Material designated as "CONFIDENTIAL ATTORNEYS' EYES ONLY" may be disclosed only to:
- (i) The Receiving Party's Outside Counsel, provided that such Outside Counsel is not involved in competitive decision-making, as defined by *U.S. Steel v. United States*, 730 F.2d 1465, 1468 n.3 (Fed. Cir. 1984), on behalf of a Party or a competitor of a Party, and such

Outside Counsel's immediate paralegals and staff, and any copying or clerical litigation support services working at the direction of such counsel, paralegals, and staff;

assist in this action, provided that disclosure is only to the extent necessary to perform such work; and provided that: (a) such expert or consultant has agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A; (b) such expert or consultant is not a current officer, director, or employee of a Party or of a competitor of a Party, nor anticipated at the time of retention to become an officer, director, or employee of a Party or of a competitor of a Party; (c) such expert or consultant is not involved in competitive decision-making, as defined by *U.S. Steel v. United States*, 730 F.2d 1465, 1468 n.3 (Fed. Cir. 1984), on behalf of a Party or a competitor of a Party; (d) such expert or consultant accesses the materials in the United States only, and does not transport them to or access them from any foreign jurisdiction (however, to avoid doubt, such expert or consultant may access reports (including drafts) that contain the materials for purposes of working on these cases while traveling temporarily outside the United States); and (e) no unresolved objections to such disclosure exist after proper notice has been given to all Parties as set forth in Section 12 below;

(iii) Witnesses at depositions or hearings in these cases and the witnesses' counsel, provided however that the disclosure shall only be made to: (1) a witness who is identified on the document as an author, addressee, or recipient of the material in question, or if there are other indicia (such as from testimony, metadata, cover emails, or other records of distribution) that the witness has previously seen or had access to the document or the information contained therein; or (2) a witness who has been designated to testify on behalf of the Producing Party on the subject matter of the material in question, provided however that

the Protected Material shown to such a witness shall be limited to Protected Material of the Producing Party;

- (iv) Court reporters, stenographers and videographers retained to record testimony taken in this action, and their staff;
 - (v) The Court, jury, and court personnel;
- (vi) Graphics, translation, design, trial consulting personnel, and/or other professional vendors, having first agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A;
- (vii) Any mediator who is assigned to hear this matter, and his or her staff, subject to their agreement to maintain confidentiality to the same degree as required by this Protective Order; and
- (viii) Any other person with the prior written consent of the Producing Party.
- (c) In addition, a Party may disclose arguments and materials derived from Discovery Material designated as "CONFIDENTIAL ATTORNEYS' EYES ONLY" to mock jurors who have signed an undertaking or agreement agreeing not to publicly disclose Protected Material and to keep any information concerning Protected Material confidential. A Party may not disclose to mock jurors any original, as-produced materials or information (including, for example, documents, deposition testimony, or interrogatory responses) produced by another Party designated as "CONFIDENTIAL ATTORNEYS' EYES ONLY."

10. <u>DISCOVERY MATERIAL DESIGNATED AS "CONFIDENTIAL – OUTSIDE ATTORNEYS' EYES ONLY - SOURCE CODE"</u>

(a) To the extent production of Source Code becomes necessary to the prosecution or defense of the cases, a Producing Party may designate Source Code as

"CONFIDENTIAL – OUTSIDE ATTORNEYS' EYES ONLY - SOURCE CODE" if it comprises or includes confidential, proprietary, and/or trade secret Source Code.

- (b) Nothing in this Order shall be construed as a representation or admission that Source Code is properly discoverable in these cases, or to obligate any Party to produce any Source Code.
- (c) Unless otherwise ordered by the Court, Discovery Material designated as "CONFIDENTIAL OUTSIDE ATTORNEYS' EYES ONLY SOURCE CODE" shall be subject to the provisions set forth in Section 11 below, and may be disclosed, subject to Section 11 below, solely to:
- (i) The Receiving Party's Outside Counsel, provided that such Outside Counsel is not involved in competitive decision-making, as defined by *U.S. Steel v. United States*, 730 F.2d 1465, 1468 n.3 (Fed. Cir. 1984), on behalf of a Party or a competitor of a Party, and such Outside Counsel's immediate paralegals and staff, and any copying or clerical litigation support services working at the direction of such counsel, paralegals, and staff;
- (ii) Any outside expert or consultant retained by the Receiving Party to assist in this action, provided that disclosure is only to the extent necessary to perform such work; and provided that: (a) such expert or consultant has agreed to be bound by the provisions of the Protective Order by signing a copy of Exhibit A; (b) such expert or consultant is not a current officer, director, or employee of a Party or of a competitor of a Party, nor anticipated at the time of retention to become an officer, director or employee of a Party or of a competitor of a Party; (c) such expert or consultant is not involved in competitive decision-making, as defined by *U.S. Steel v. United States*, 730 F.2d 1465, 1468 n.3 (Fed. Cir. 1984), on behalf of a Party or a competitor of a Party; (d) such expert or consultant accesses the materials in the United States only, and does not

transport them to or access them from any foreign jurisdiction; and (e) no unresolved objections to such disclosure exist after proper notice has been given to all Parties as set forth in Section 12 below;

- (iii) Witnesses at depositions or hearings in these cases and the witnesses' counsel, provided however that the disclosure shall only be made to: (1) a witness who is identified on the material as an author, addressee, or recipient of the material, or if there are indicia (such as from testimony, metadata, emails, or other records of distribution) that the witness has seen or had access to the materials previously; or (2) a witness who has been designated to testify on behalf of the Producing Party on the subject matter of the material in question, provided however that the Protected Material shown to such a witness shall be limited to Protected Material of the Producing Party;
- (iv) Court reporters, stenographers and videographers retained to record testimony taken in this action, and their staff;
 - (v) The Court, jury, and court personnel;
- (vi) Any mediator who is assigned to hear this matter, and his or her staff, subject to their agreement to maintain confidentiality to the same degree as required by this Protective Order; and
- (vii) Any other person with the prior written consent of the Producing Party.

11. <u>DISCLOSURE AND REVIEW OF SOURCE CODE</u>

(a) Any Source Code that is produced by Plaintiff will be made available for inspection at the San Francisco office of its outside counsel, Desmarais LLP, or any other location mutually agreed by the Parties. Any Source Code that is produced by Masimo will be made

available for inspection at the Orange County office of their outside counsel, Knobbe Martens Olsen & Bear LLP, or any other location mutually agreed by the Parties. Source Code will be made available for inspection between the hours of 8 a.m. and 6 p.m. on business days (i.e., weekdays that are not Federal holidays), although the Parties will be reasonable in accommodating reasonable requests to conduct inspections at other times.

- (b) Prior to the first inspection of any requested Source Code, the Receiving Party shall provide ten (10) days' notice of its intent to review the Source Code that has been made available by the Producing Party and, if known, the specific Source Code the Receiving Party intends to inspect. The Receiving Party shall provide seven (7) days' notice prior to any additional inspections.
- (c) Source Code that is designated "CONFIDENTIAL OUTSIDE ATTORNEYS' EYES ONLY SOURCE CODE" shall be produced for inspection and review subject to the following provisions, unless otherwise agreed by the Producing Party:
- the Receiving Party's Outside Counsel and/or experts in a secure room on a secured computer without Internet access or network access to other computers and on which all access ports have been disabled (except for one printer port), as necessary and appropriate to prevent and protect against any unauthorized copying, transmission, removal or other transfer of any Source Code outside or away from the computer on which the Source Code is provided for inspection (the "Source Code Computer" in the "Source Code Review Room"). The Producing Party shall install tools that are sufficient for viewing and searching the code produced, on the platform produced, if such tools exist and are presently used in the ordinary course of the Producing Party's business. The Receiving Party's Outside Counsel and/or experts may request that commercially available

software tools for viewing and searching Source Code be installed on the secured computer, provided, however, that (a) the Receiving Party possesses an appropriate license to such software tools; (b) the Producing Party approves such software tools (approvals will not be unreasonably denied); and (c) such other software tools are reasonably necessary for the Receiving Party to perform its review of the Source Code consistent with all of the protections herein. The Receiving Party must provide the Producing Party with the CD or DVD or other media containing such licensed software tool(s) at least seven (7) days in advance of the date upon which the Receiving Party wishes to have the additional software tools available for use on the Source Code Computer.

- (ii) No recordable media or recordable devices, including without limitation sound recorders, computers, cellular telephones, peripheral equipment, cameras, CDs, DVDs, or drives of any kind, shall be permitted into the Source Code Review Room.
- (iii) The Receiving Party's Outside Counsel and/or experts shall be entitled to take notes relating to the Source Code but may not copy the Source Code into the notes and may not take such notes electronically on the Source Code Computer itself or any other computer.
- (iv) The Producing Party may visually monitor the activities of the Receiving Party's representatives during any Source Code review, but only to ensure that no unauthorized electronic records of the Source Code and no information concerning the Source Code are being created or transmitted in any way.
- (v) No copies of all or any portion of the Source Code may leave the room in which the Source Code is inspected except as otherwise provided herein. Further, no other written or electronic record of the Source Code is permitted except as otherwise provided herein. The Producing Party shall make available a laser printer with commercially reasonable

printing speeds for on-site printing during inspection of the Source Code. The Receiving Party may print limited portions of the Source Code only when necessary to prepare court filings or pleadings or other papers (including a testifying expert's expert report). The Receiving Party may print the Source Code in 12-point font and with information necessary to later identify that Source Code, such as, but not limited to, a header or footer, that identifies the file name and directory path. Any printed portion that consists of more than fifteen (15) pages of a continuous block of Source Code shall be presumed to be excessive, and the burden shall be on the Receiving Party to demonstrate the need for such a printed copy. The Receiving Party may print out no more than 200 pages total without prior agreement from the Producing Party or order of the Court. The Receiving Party shall not print Source Code in order to review blocks of Source Code elsewhere in the first instance, i.e., as an alternative to reviewing that Source Code electronically on the Source Code Computer, as the Parties acknowledge and agree that the purpose of the protections herein would be frustrated by printing portions of code for review and analysis elsewhere, and that printing is permitted only when necessary to prepare court filings or pleadings or other papers (including a testifying expert's expert report). Upon printing any such portions of Source Code, the printed pages shall be collected by the Producing Party. The Producing Party shall Bates number, copy, and label "CONFIDENTIAL - OUTSIDE ATTORNEYS" EYES ONLY -SOURCE CODE" any pages printed by the Receiving Party. Within seven (7) days, the Producing Party shall either (i) provide one copy set of such pages to the Receiving Party or (ii) inform the Requesting Party that it objects that the printed portions are excessive and/or not done for a permitted purpose. If, after meeting and conferring, the Producing Party and the Receiving Party cannot resolve the objection, the Receiving Party shall be entitled to seek a Court resolution of whether the printed Source Code in question is narrowly tailored and was printed for a permitted purpose. The

burden shall be on the Receiving Party to demonstrate that such printed portions are no more than is reasonably necessary for a permitted purpose and not merely printed for the purposes of review and analysis elsewhere. The printed pages shall constitute part of the Source Code produced by the Producing Party in these cases.

(vi) All persons who will review a Producing Party's Source Code on behalf of a Receiving Party, including members of a Receiving Party's outside law firm, shall be identified in writing to the Producing Party at least five (5) days in advance of the first time that such person reviews such Source Code. Such identification shall be in addition to any other disclosure required under this Order. All persons viewing Source Code shall sign on each day they view Source Code a log that will include the names of persons who enter the locked room to view the Source Code and when they enter and depart. The Producing Party shall be entitled to a copy of the log upon one (1) day's advance notice to the Receiving Party.

(vii) Unless otherwise agreed in advance by the Parties in writing, following each day on which inspection is done under this Order, the Receiving Party's Outside Counsel and/or experts shall remove all notes, documents, and all other materials from the Source Code Review Room. The Producing Party shall not be responsible for any items left in the room following each inspection session, and the Receiving Party shall have no expectation of confidentiality for any items left in the room following each inspection session without a prior agreement to that effect. Proper identification of all authorized persons shall be provided prior to any access to the secure room or the computer containing Source Code. Proper identification requires showing, at a minimum, a photo identification card sanctioned by the government of any State of the United States, by the government of the United States, or by the nation state of the authorized person's current citizenship. Access to the secure room or the Source Code Computer

may be denied, at the discretion of the supplier, to any individual who fails to provide proper identification.

- (viii) Other than as provided above, the Receiving Party will not copy, remove, or otherwise transfer any Source Code from the Source Code Computer including, without limitation, copying, removing, or transferring the Source Code onto any recordable media or recordable device. The Receiving Party will not transmit any Source Code in any way from the Producing Party's facilities or the offices of its Outside Counsel of record.
- (ix) The Receiving Party's Outside Counsel of record may make no more than three (3) additional paper copies of any portions of the Source Code received from a Producing Party pursuant to Section 11(c)(v), not including copies attached to court filings or used at depositions, and shall maintain a log of all paper copies of the Source Code. The log shall include the names of the reviewers and/or recipients of paper copies and locations where the paper copies are stored. Upon one (1) day's advance notice to the Receiving Party by the Producing Party, the Receiving Party shall provide a copy of this log to the Producing Party.
- (x) The Receiving Party's Outside Counsel of record and any person receiving a copy of any Source Code shall maintain and store any paper copies of the Source Code at their offices in a manner that prevents duplication of or unauthorized access to the Source Code, including, without limitation, storing the Source Code in a locked room or cabinet at all times when it is not in use. No more than a total of fifteen (15) individuals identified by the Receiving Party shall have access to the printed portions of Source Code (except insofar as such code appears in any court filing or expert report).
- (xi) For depositions, the Receiving Party shall not bring copies of any printed Source Code. Rather, at least seven (7) days before the date of the deposition, the Receiving

Party shall notify the Producing Party about the specific portions of Source Code it wishes to use at the deposition, and the Producing Party shall bring printed copies of those portions to the deposition for use by the Receiving Party. The Producing Party shall also accommodate reasonable requests from the Receiving Party to make a Source Code Computer available at the deposition for use at the deposition. Copies of Source Code that are marked as deposition exhibits shall not be provided to the Court Reporter or attached to deposition transcripts; rather, the deposition record will identify the exhibit by its production numbers. All paper copies of Source Code brought to the deposition shall remain with the Producing Counsel's Outside Counsel for secure destruction in a timely manner following the deposition.

(xii) Except as provided in this section, absent express written permission from the Producing Party, the Receiving Party may not create electronic images, or any other images, or make electronic copies, of the Source Code from any paper copy of Source Code for use in any manner (including by way of example only, the Receiving Party may not scan the Source Code to a PDF or photograph the code). Images or copies of Source Code shall not be included in correspondence between the Parties (references to production numbers shall be used instead), and shall be omitted from pleadings and other papers whenever possible. If a Party reasonably believes that it needs to submit a portion of Source Code as part of a filing with the Court, the Parties shall meet and confer as to how to make such a filing while protecting the confidentiality of the Source Code and such Source Code will not be filed absent agreement from the Producing Party that the confidentiality protections will be adequate. If a Producing Party agrees to produce an electronic copy of all or any portion of its Source Code or provide written permission to the Receiving Party that an electronic or any other copy needs to be made for a Court filing, access to the Receiving Party's submission, communication, and/or disclosure of electronic files or other materials

containing any portion of Source Code (paper or electronic) shall at all times be limited solely to individuals who are expressly authorized to view Source Code under the provisions of this Order. Where the Producing Party has provided the express written permission required under this provision for a Receiving Party to create electronic copies of Source Code, the Receiving Party shall maintain a log of all such electronic copies of any portion of Source Code in its possession or in the possession of its retained consultants, including the names of the reviewers and/or recipients of any such electronic copies, and the locations and manner in which the electronic copies are stored. Additionally, any such electronic copies must be labeled "CONFIDENTIAL - ATTORNEYS' EYES ONLY - SOURCE CODE" as provided for in this Order.

12. NOTICE OF DISCLOSURE

- (a) Prior to disclosing any Protected Material to any person described in Sections 8(b)(iii), 9(b)(ii), or 10(c)(ii) (referenced below as "Person"), the Party seeking to disclose such information shall provide the Producing Party with written notice that includes:
 - (i) the name of the Person;
 - (ii) an up-to-date curriculum vitae of the Person;
 - (iii) the present employer and title of the Person;
- (iv) an identification of all of the Person's past and current employment and consulting relationships in the past five years, including direct relationships and relationships through entities owned or controlled by the Person, including but not limited to an identification of any individual or entity with or for whom the person is employed or to whom the person provides consulting services relating to the design, development, operation, or patenting of technologies relating to non-invasive monitoring and/or consumer wearables (generally or as described in any patent in suit), or relating to the acquisition of intellectual property assets relating

to non-invasive monitoring and/or consumer wearables (generally or as described in any patent in suit);

- (v) an identification of all pending patent applications on which the Person is named as an inventor, in which the Person has any ownership interest, or as to which the Person has had or anticipates in the future any involvement in advising on, consulting on, preparing, prosecuting, drafting, editing, amending, or otherwise affecting the scope of the claims; and
- (vi) a list of the cases in which the Person has testified at deposition or trial within the last five (5) years.

Further, the Party seeking to disclose Protected Material shall provide such other information regarding the Person's professional activities reasonably requested by the Producing Party for it to evaluate whether good cause exists to object to the disclosure of Protected Material to the outside expert or consultant.

Party or Parties may object in writing to the Person for good cause. In the absence of an objection at the end of the ten (10) day period, the Person shall be deemed approved under this Protective Order. There shall be no disclosure of Protected Material to the Person prior to expiration of this ten (10) day period. If the Producing Party objects to disclosure to the Person within such ten (10) day period, the Parties shall meet and confer via telephone or in person within four (4) days following the objection and attempt in good faith to resolve the dispute on an informal basis. If the dispute is not resolved, the Party objecting to the disclosure will have four (4) days from the date of the meet and confer to seek relief from the Court and shall have the burden of proving the need for a protective order. If relief is not sought from the Court within that time, the objection

shall be deemed withdrawn. If relief is sought, designated materials shall not be disclosed to the Person in question until the Court resolves the objection.

- (c) For purposes of this section, "good cause" shall include an objectively reasonable concern that the Person will, advertently or inadvertently, use or disclose Discovery Material in a way or ways that are inconsistent with the provisions contained in this Order.
- (d) Prior to receiving any Protected Material under this Order, the Person must execute a copy of the "Agreement to Be Bound by Protective Order" (Exhibit A hereto) and serve it on all Parties.
- (e) An initial failure to object to a Person under this Section 12 shall not preclude the nonobjecting Party from later objecting to continued access by that Person for good cause. If an objection is made, the Parties shall meet and confer via telephone or in person within seven (7) days following the objection and attempt in good faith to resolve the dispute informally. If the dispute is not resolved, the Party objecting to the disclosure will have seven (7) days from the date of the meet and confer to seek relief from the Court. The designated Person may continue to have access to information that was provided to such Person prior to the date of the objection. If a later objection is made, no further Protected Material shall be disclosed to the Person until the Court resolves the matter or the Producing Party withdraws its objection. Notwithstanding the foregoing, if the Producing Party fails to move for a protective order within seven (7) business days after the meet and confer, further Protected Material may thereafter be provided to the Person.

13. CHALLENGING DESIGNATIONS OF PROTECTED MATERIAL

(a) A Party shall not be obligated to challenge the propriety of any designation of Discovery Material under this Order at the time the designation is made, and a failure to do so shall not preclude a subsequent challenge thereto.

- (b) Any challenge to a designation of Discovery Material under this Order shall be written, shall be served on Outside Counsel for the Producing Party, shall particularly identify the documents or information that the Receiving Party contends should be differently designated, and shall state the grounds for the objection. Thereafter, further protection of such material shall be resolved in accordance with the following procedures:
- (i) The objecting Party shall have the burden of conferring either in person, in writing, or by telephone with the Producing Party claiming protection (as well as any other interested party) in a good faith effort to resolve the dispute. The Producing Party shall have the burden of justifying the disputed designation;
- (ii) Failing agreement, the Receiving Party may bring a request or motion to the Court for a ruling that the Discovery Material in question is not entitled to the status and protection of the Producing Party's designation. The Parties' entry into this Order shall not preclude or prejudice either Party from arguing for or against any designation, establish any presumption that a particular designation is valid, or alter the burden of proof that would otherwise apply in a dispute over discovery or disclosure of information;
- (iii) Notwithstanding any challenge to a designation, the Discovery Material in question shall continue to be treated as designated under this Order until one of the following occurs: (a) the Party who designated the Discovery Material in question withdraws such designation in writing; or (b) the Court rules that the Discovery Material in question is not entitled to the designation.

14. **DATA SECURITY**

(a) The Receiving Party shall implement an information security management system ("ISMS") to safeguard Protected Materials, including reasonable and appropriate

administrative, physical, and technical safeguards, and network security and encryption technologies governed by written policies and procedures, which shall comply with at least one of the following standards: (a) the International Organization for Standardization's 27001 standard; (b) the National Institute of Standards and Technology's (NIST) 800-53 standard; (c) the Center for Internet Security's Critical Security Controls, Version 8; or (d) the most recently published version of another widely recognized industry or government cybersecurity framework. The Parties shall implement encryption of all Protected Materials in transit outside of network(s) covered by the Party's ISMS (and at rest, where reasonably practical). Moreover, the Parties agree not to access Protected Materials from public computers.

- disclosure of Protected Materials or devices containing Protected Materials ("Data Breach"), the Receiving Party shall promptly, and in no case later than 48 hours after learning of the Data Breach, notify the Producing Party in writing and fully cooperate with the Producing Party as may be reasonably necessary to (a) determine the source, extent, or methodology of such Data Breach, (b) recover or protect Protected Materials, and/or (c) to satisfy the Producing Party's legal, contractual, or other obligations. For the avoidance of doubt, notification obligations under this section arise when the Receiving Party both (a) learns of a Data Breach, and (b) learns that any of the Producing Party's Protected Materials are potentially subject to the Data Breach. The notification obligations set forth in this section do not run from the time the Data Breach itself.
- (c) If the Receiving Party is aware of a Data Breach, the Parties shall meet and confer in good faith regarding any adjustments that should be made to the discovery process and discovery schedule in these cases, potentially including but not limited to (1) additional security measures to protect Discovery Material; (2) a stay or extension of discovery pending investigation

of a Data Breach and/or implementation of additional security measures; and (3) a sworn assurance that Discovery Material will be handled in the future only by entities not impacted by the Data Breach. In the event of a Data Breach affecting Protected Material of the Designating Party, at the Designating Party's request, the Receiving Party within 10 business days shall provide a copy of its most recent ISMS policies and procedures that relate to the safeguarding of Protected Materials and that preceded the Data Breach. Further, the Receiving Party shall submit to reasonable discovery concerning the Data Breach.

15. SUBPOENAS OR COURT ORDERS

(a) If at any time Protected Material is subpoenaed by any court, arbitral, administrative, or legislative body, the Party to whom the subpoena or other request is directed shall immediately give prompt written notice thereof to every Party who has produced such Discovery Material and to its counsel and shall provide each such Party with an opportunity to move for a protective order regarding the production of Protected Materials implicated by the subpoena. The Producing Party must also notify in writing the party who caused the subpoena or order to issue in the other litigation that some or all of the material covered by the subpoena or order is subject to this Protective Order, and include a copy of this Protective Order. The parties agree to work together to allow the Producing Party to seek a protective order, after the filing of which the Party served with the subpoena or court order shall not produce any information designated in this action as "CONFIDENTIAL – ATTORNEYS EYES ONLY" or "CONFIDENTIAL – ATTORNEYS EYES ONLY – SOURCE CODE" before a determination on the protective order by the court from which the subpoena or order issued, unless the Party has obtained the Producing Party's permission.

16. **FILING PROTECTED MATERIAL**

- (a) Absent written permission from the Producing Party or a court Order secured after appropriate notice to all interested persons, a Receiving Party may not file or disclose in the public record any Protected Material.
- (b) Any Party is authorized under District of Delaware Local Rule 5.1.3 to file under seal with the Court any brief, document or materials that are designated as Protected Material under this Order. However, nothing in this section shall in any way limit or detract from this Order's requirements as to Source Code.

17. INADVERTENT DISCLOSURE OF PRIVILEGED MATERIAL

(a) Pursuant to Federal Rule of Evidence 502(d) and (e), the inadvertent production by a Party of Discovery Material subject to the attorney-client privilege, work-product protection, or any other applicable privilege or protection, despite the Producing Party's reasonable efforts to prescreen such Discovery Material prior to production, will not waive the applicable privilege and/or protection in any other federal or state proceeding if a request for return of such inadvertently produced Discovery Material is made promptly after the Producing Party learns of its inadvertent production. For example, the mere production of a privileged or work product protected document in this case as part of a production is not itself a waiver. Nothing in this Order shall be interpreted to require disclosure of irrelevant information or relevant information protected by the attorney-client privilege, work product doctrine, or any other applicable privilege or immunity. The parties do not waive any objections as to the production, discoverability, admissibility, or confidentiality of documents and ESI. Moreover, nothing in this Order shall be interpreted to require disclosure of information subject to privacy protections as set forth in law or

regulation, including information that may need to be produced from outside of the United States and/or may be subject to foreign laws.

- (b) Upon a request from any Producing Party who has inadvertently produced Discovery Material that it believes is privileged and/or protected, each Receiving Party shall immediately return such Protected Material or Discovery Material and all copies to the Producing Party, except for any pages containing privileged markings by the Receiving Party which shall instead be destroyed and certified as such by the Receiving Party to the Producing Party.
- (c) Nothing herein shall prevent the Receiving Party from preparing a record for its own use containing the date, author, addresses, and topic of the inadvertently produced Discovery Material and such other information as is reasonably necessary to identify the Discovery Material and describe its nature to the Court in any motion to compel production of the Discovery Material.

18. <u>INADVERTENT FAILURE TO DESIGNATE PROPERLY</u>

- Material as Protected Material with one of the designations provided for under this Order shall not waive any such designation provided that the Producing Party notifies all Receiving Parties that such Discovery Material is protected under one of the categories of this Order within ten (10) days of the Producing Party learning of the inadvertent failure to designate. The Producing Party shall reproduce the Protected Material with the correct confidentiality designation within five (5) days upon its notification to the Receiving Parties. Upon receiving the Protected Material with the correct confidentiality designation, the Receiving Parties shall return or securely destroy, at the Producing Party's option, all Discovery Material that was not designated properly.
- (b) A Receiving Party shall not be in breach of this Order for any use of such Discovery Material before the Receiving Party receives such notice that such Discovery Material

is protected under one of the categories of this Order, unless an objectively reasonable person would have realized that the Discovery Material should have been appropriately designated with a confidentiality designation under this Order. Once a Receiving Party has received notification of the correct confidentiality designation for the Protected Material with the correct confidentiality designation, the Receiving Party shall treat such Discovery Material (subject to the exception in Section 18(c) below) at the appropriately designated level pursuant to the terms of this Order.

(c) Notwithstanding the above, a subsequent designation of "CONFIDENTIAL," "CONFIDENTIAL – ATTORNEYS' EYES ONLY" or "CONFIDENTIAL – ATTORNEYS' EYES ONLY – SOURCE CODE" shall apply on a going forward basis and shall not disqualify anyone who reviewed "CONFIDENTIAL," "CONFIDENTIAL – ATTORNEYS' EYES ONLY" or "CONFIDENTIAL – ATTORNEYS' EYES ONLY – SOURCE CODE" materials while the materials were not marked "CONFIDENTIAL – ATTORNEYS' EYES ONLY" or "CONFIDENTIAL – ATTORNEYS' EYES ONLY – SOURCE CODE" from engaging in the activities set forth in Section 6(b).

19. <u>INADVERTENT DISCLOSURE NOT AUTHORIZED BY ORDER</u>

(a) In the event of a disclosure of any Discovery Material pursuant to this Order to any person or persons not authorized to receive such disclosure under this Protective Order, the Party responsible for having made such disclosure, and each Party with knowledge thereof, shall immediately notify counsel for the Producing Party whose Discovery Material has been disclosed and provide to such counsel all known relevant information concerning the nature and circumstances of the disclosure. The responsible disclosing Party shall also promptly take all reasonable measures to retrieve the improperly disclosed Discovery Material and to ensure that no further or greater unauthorized disclosure and/or use thereof is made.

(b) Unauthorized or inadvertent disclosure does not change the status of Discovery Material or waive the right to hold the disclosed document or information as Protected.

20. **FINAL DISPOSITION**

- (a) Not later than ninety (90) days after the Final Disposition of these cases, each Party shall return all Discovery Material of a Producing Party to the respective Outside Counsel of the Producing Party or destroy such Material, at the option of the Producing Party. For purposes of this Order, "Final Disposition" occurs after an order, mandate, or dismissal finally terminating these cases with prejudice, including all appeals.
- (b) All Parties that have received any such Discovery Material shall certify in writing that all such materials have been returned to the respective Outside Counsel of the Producing Party or destroyed. Notwithstanding the provisions for return of Discovery Material, Outside Counsel may retain one set of pleadings, correspondence and attorney and consultant work product (but not document productions) for archival purposes, but must return any pleadings, correspondence, and consultant work product that contain Source Code.

21. MISCELLANEOUS

- (a) <u>Right to Further Relief.</u> Nothing in this Order abridges the right of any person to seek its modification by the Court in the future. By stipulating to this Order, the Parties do not waive the right to argue that certain material may require additional or different confidentiality protections than those set forth herein.
- (b) <u>Termination of Matters and Retention of Jurisdiction</u>. The Parties agree that the terms of this Protective Order shall survive and remain in effect after the Final Determination of the above-captioned matters. The Court shall retain jurisdiction after Final Determination of these matters to hear and resolve any disputes arising out of this Protective Order.

- (c) <u>Successors</u>. This Order shall be binding upon the Parties hereto, their successors, and anyone, including law firms, who obtains access to Protected Material.
- (d) Right to Assert Other Objections. By stipulating to the entry of this Protective Order, no Party waives any right it otherwise would have to object to disclosing or producing any information or item. Similarly, no Party waives any right to object on any ground to use in evidence of any of the material covered by this Protective Order. This Order shall not constitute a waiver of the right of any Party to claim in these cases or otherwise that any Discovery Material, or any portion thereof, is privileged or otherwise non-discoverable, or is not admissible in evidence in these cases or any other proceeding.
- (e) <u>Modification by Court</u>. This Order is subject to further court order based upon public policy or other considerations, and the Court may modify this Order *sua sponte* in the interests of justice. The United States District Court for the District of Delaware is responsible for the interpretation and enforcement of this Order. All disputes concerning Protected Material, however designated, produced under the protection of this Order shall be resolved by the United States District Court for the District of Delaware.

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Attorneys for Defendants Masimo Corporation and Sound United, LLC

Attorneys for Plaintiff Apple Inc.

Dated: June 14, 2023

IT IS SO ORDERED this 16th day of June, 2023.

The Honorable Jennifer L. Hall

United States District Court Magistrate Judge

EXHIBIT A

| I, | , acknowledge and declare that I have received a |
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| copy of the Prote | ctive Order ("Order") in Apple Inc. v. Masimo Corp. et al., United States |
| District Court, D | vistrict of Delaware, C.A. Nos. 22-1377-MN-JLH and 22-1378-MN-JLH. |
| Having read and | understood the terms of the Order, I agree to be bound by the terms of the |
| Order and consent | to the jurisdiction of said Court for the purpose of any proceeding to enforce |
| the terms of the O | rder. |
| Name of i | ndividual: |
| Present occ | cupation/job description: |
| | |
| | ompany or Firm: |
| Address: | |
| Dated: | |
| | [Signature] |

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